

5_3_Magic_BLAST_contigs_specific

July 25, 2021

Summary of magicblast

```
[1]: import numpy as np
import pandas as pd
from matplotlib import pyplot as plt
import seaborn as sns
from pandas.plotting import scatter_matrix
import pathlib
import warnings
warnings.filterwarnings('ignore')
```

```
[2]: from IPython.core.display import display, HTML
display(HTML("<style>.container { width:95% !important; }</style>"))
```

<IPython.core.display.HTML object>

```
[3]: PROJECT_CODE='PRJNA573298'
BASE_PATH = f'/mnt/1TB_0/Data/Assembly/{PROJECT_CODE}/'

dbname='nt'
kmer='k141'

#magicblast on contigs
magic_blast_sam_tail=f'_{dbname}_final_contigs_magicBLAST.sam'
f_contigs_file_tail=f'_{dbname}_magic_blast_asc_contigs.txt'
GIS,ACCESSIONS,TITLES=None,None,None
```

0.0.1 All SRA's

```
[4]: sra_list=['SRR10168373','SRR10168374',\
              'SRR10168375','SRR10168376',\
              'SRR10168377','SRR10168378',\
              'SRR10168379','SRR10168380',\
              'SRR10168381','SRR10168382',\
              'SRR10168383','SRR10168384',\
              'SRR10168385','SRR10168386',\
              'SRR10168387','SRR10168388',\
              'SRR10168389','SRR10168390',\
```

```
'SRR10168391','SRR10168392','SRR10168393']
```

```
[5]: def read_gi_accession_title(gi_asc_file):  
    gis=[]  
    accessions=[]  
    titles=[]  
    with open(gi_asc_file, 'r') as infile:  
        data = infile.readlines()  
        for i in data:  
            output=i.split(' ',2)  
            gis.append(output[0])  
            accessions.append(output[1])  
            titles.append(output[2])  
    return gis, accessions, titles
```

```
[6]: def set_accessions():  
    global GIS  
    global ACCESSIONS  
    global TITLES  
    GIS,ACCESSIONS,TITLES=read_gi_accession_title('/mnt/1TB_ssd/Data/BLAST/nt.  
↳gi_taxid.tsv')
```

0.0.2 Stats

```
[7]: def get_asc_descr_count(sra):  
    accessions=[]  
    descriptions=[]  
    counts=[]  
    path = BASE_PATH+sra+'/magic_blast/'  
    with open(path+f'{sra}_{dbname}_{kmer}_magicBLAST_summary.txt', 'r') as f:  
        lines = [line.rstrip('\n') for line in f]  
        total=len(lines)-1  
        for line in lines:  
            if not 'database:' in line: #ignore header  
                asc=line.split(None, 1)[0]  
                title=line.split(None, 1)[1].split(', count:')[0]  
                count=line.split(None, 1)[1].split(', count:')[1]  
                accessions.append(asc)  
                descriptions.append(title)  
                counts.append(int(count))  
    return accessions, descriptions, counts, total
```

```
[8]: def get_indexes(substr, str_list):  
    index_list = []  
    i = 0  
    for e in str_list:  
        if substr in e.lower():
```

```

        index_list.append(i)
    i +=1
    return index_list

```

```

[9]: for i,sra in enumerate(sra_list):
      accessions, descriptions, counts, total=get_asc_descr_count(sra)
      print(f'{sra}, matched contigs: {sum(counts)}')
      #for d,c in zip(descriptions, counts):
      #    print(f'{d}: {c}')
      #print('\n')

```

```

SRR10168373, matched contigs: 4348
SRR10168374, matched contigs: 4669
SRR10168375, matched contigs: 43799
SRR10168376, matched contigs: 23385
SRR10168377, matched contigs: 19599
SRR10168378, matched contigs: 14969
SRR10168379, matched contigs: 45840
SRR10168380, matched contigs: 33249
SRR10168381, matched contigs: 18395
SRR10168382, matched contigs: 3050
SRR10168383, matched contigs: 4954
SRR10168384, matched contigs: 15128
SRR10168385, matched contigs: 4704
SRR10168386, matched contigs: 2146
SRR10168387, matched contigs: 9448
SRR10168388, matched contigs: 4383
SRR10168389, matched contigs: 4398
SRR10168390, matched contigs: 8835
SRR10168391, matched contigs: 18251
SRR10168392, matched contigs: 35405
SRR10168393, matched contigs: 18335

```

```

[10]: def get_desc_count(qstring, descriptions, counts, lowercase=True):
      qd=[]
      qc=0
      for d,c in zip(descriptions, counts):
          if lowercase:
              if qstring.lower() in d.lower():
                  qd.append(d)
                  qc=qc+int(c)
          else:
              if qstring in d:
                  qd.append(d)
                  qc=qc+int(c)
      #print(f'qstring: {qstring}, descriptions: {len(descriptions)}, qd: {qd},
      ↪ counts: {len(counts)}, qc: {qc}')

```

```
return qd, qc
```

```
[11]: def get_desc_count_without(qstring, nqstring, descriptions, counts,
↳ lowercase=True):
    qd=[]
    qc=0
    for d,c in zip(descriptions, counts):
        if lowercase:
            if qstring.lower() in d.lower() and nqstring.lower() not in d.
↳ lower():
                qd.append(d)
                qc=qc+int(c)
        else:
            if qstring in d:
                qd.append(d)
                qc=qc+int(c)
        #print(f'qstring: {qstring}, descriptions: {len(descriptions)}, qd: {qd},
↳ counts: {len(counts)}, qc: {qc}')
    return qd, qc
```

```
[12]: def get_descr(sra):
    accessions, descriptions, counts, total=get_asc_descr_count(sra)
    #print(f'sra: {sra}, accessions: {len(accessions)}, descriptions:
↳ {len(descriptions)}, counts: {len(counts)}, total: {total}')
    if total>0:
        human,humanc = get_desc_count('human', descriptions, counts)
        homo_sapiens,homo_sapiensc = get_desc_count('homo sapiens',
↳ descriptions, counts)
        h_sapiens,h_sapiensc = get_desc_count('h.sapiens', descriptions, counts)
        human_contigs = human+homo_sapiens+h_sapiens
        human_counts = humanc+homo_sapiensc+h_sapiensc
        pangolin, pangolinc = get_desc_count('manis javanica', descriptions,
↳ counts)
        pangolin_p, pangolin_pc = get_desc_count('manis pentadactyla',
↳ descriptions, counts)
        pangolin=pangolin+pangolin_p
        pangolin_counts=pangolinc+pangolin_pc
        mouse,mousec = get_desc_count('mus musculus', descriptions, counts)
        vector,vectorc= get_desc_count('vector', descriptions, counts)
        pig,pigc = get_desc_count('sus scrofa', descriptions, counts)
        cat,catc = get_desc_count('felis catus', descriptions, counts)
        tiger,tigerc = get_desc_count('panthera tigris', descriptions, counts)
        dog,dogc = get_desc_count('canis lupus', descriptions, counts)
        virus,virusc = get_desc_count_without('virus', 'retrovirus',
↳ descriptions, counts)
```

```

mulatta,mulattac = get_desc_count('mulatta', descriptions, counts)
troglodytes,troglodytesc = get_desc_count('troglodytes', descriptions,
↪counts)
pongo,pongoc =get_desc_count('pongo', descriptions, counts)
papio,papioc = get_desc_count('papio', descriptions, counts)
mandrillus,mandrillusc =get_desc_count('mandrillus', descriptions,
↪counts)
cercocebus,cercocebusc =get_desc_count('cercocebus', descriptions,
↪counts)
gelada,geladac =get_desc_count('gelada', descriptions, counts)
monkey = mulatta+troglodytes+pongo+papio+mandrillus+cercocebus+gelada
monkey_counts =
↪mulattac+troglodytesc+pongoc+papioc+mandrillusc+cercocebusc+geladac

mustela,mustelac= get_desc_count('mustela', descriptions, counts)

pipistrellus,pipistrellusc =get_desc_count('pipistrellus',
↪descriptions, counts)
rhinolophus,rhinolophusc = get_desc_count('rhinolophus', descriptions,
↪counts)
pteropus,pteropusc = get_desc_count('pteropus', descriptions, counts)
myotis,myotisc = get_desc_count('myotis', descriptions, counts)
bat = pipistrellus + rhinolophus+pteropus+myotis
bat_counts = pipistrellusc + rhinolophusc+pteropusc+myotisc

mycoplasma,mycoplasmac = get_desc_count('mycoplasma', descriptions,
↪counts)

lst = ['human', 'monkey', 'pangolin', 'mouse',
       'pig', 'cat', 'tiger', 'dog', 'bat', 'virus',
↪'vector','mycoplasma','mustela']
lengths = [human_counts, monkey_counts, pangolin_counts, mousec,
           pigc, catc, tigerc, dogc, bat_counts, virusc, vectorc,
↪mycoplasmac,mustelac]
final_contigs=BASE_PATH+sra+'/megahit_default/final.contigs.fa'
with open(final_contigs) as final_contigs_file:
    total_contigs=sum(1 for _ in final_contigs_file)
res = [int(i) for i in counts]
total_contigs_matched=sum(res)
sra_l=[sra]*len(lst)
fractions_matched = [human_counts/total_contigs_matched, monkey_counts/
↪total_contigs_matched, pangolin_counts/total_contigs_matched, mousec/
↪total_contigs_matched,

```

```

        pigc/total_contigs_matched, catc/total_contigs_matched, tiger/
↪total_contigs_matched, dogc/total_contigs_matched, bat_counts/
↪total_contigs_matched,
        virusc/total_contigs_matched, vectorc/
↪total_contigs_matched, mycoplasmac/total_contigs_matched, mustelac/
↪total_contigs_matched]
    pct_matched = [round(i * 100,2) for i in fractions_matched]
    df = pd.DataFrame(list(zip(sra_l, lst, lengths, pct_matched)),
        columns=['SRA', 'Name', 'count', 'pct_matched'])
    return df
return None

```

```

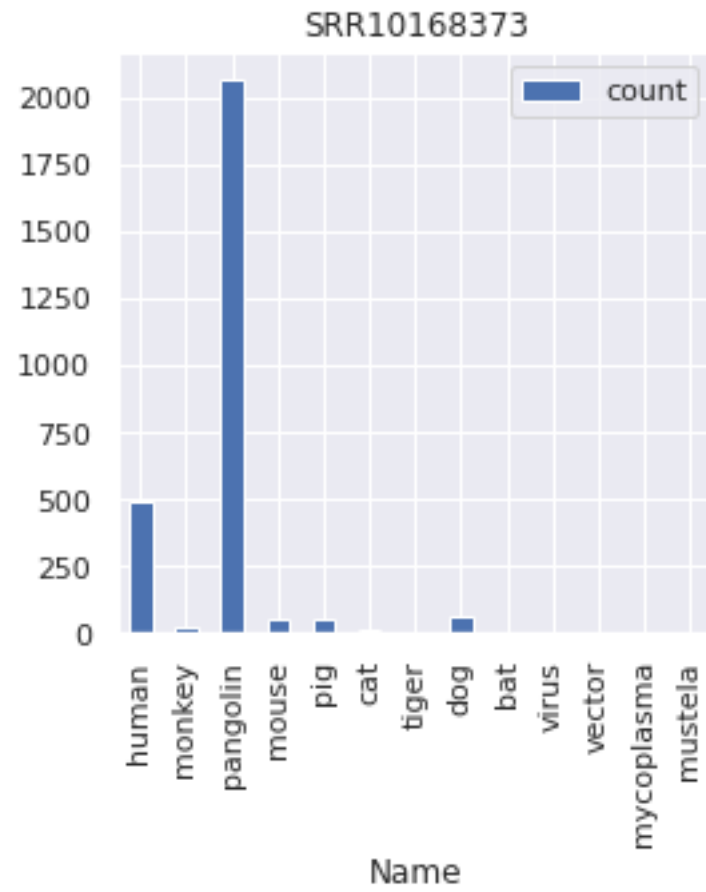
[13]: sns.set(rc={"figure.figsize":(4, 4)})
def plot_df(df, sra):
    ax=df.plot(x='Name', y='count', kind='bar')
    ax.set_title(sra, fontsize=12)
    #ax.set_yscale('log')
    #ax.set_ylim([0,df['count'].max()+10])
    ax.set_ylim(bottom=0)
    fig = plt.gcf()
    fig.savefig(BASE_PATH+sra+'/magic_blast/
↪'+f'{sra}_{dbname}_{kmer}_magicBLAST_species.png', bbox_inches="tight")

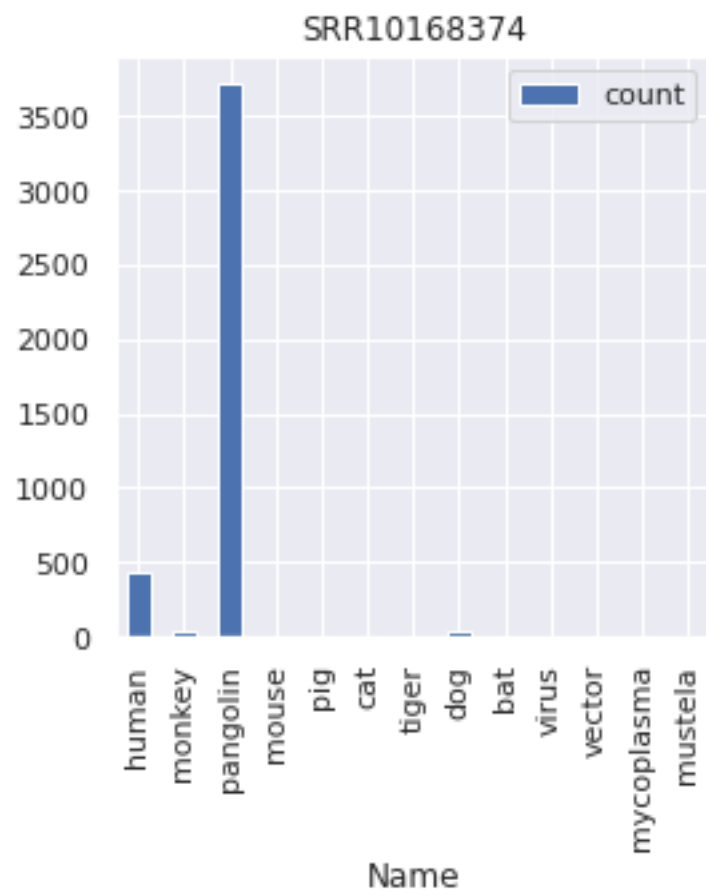
```

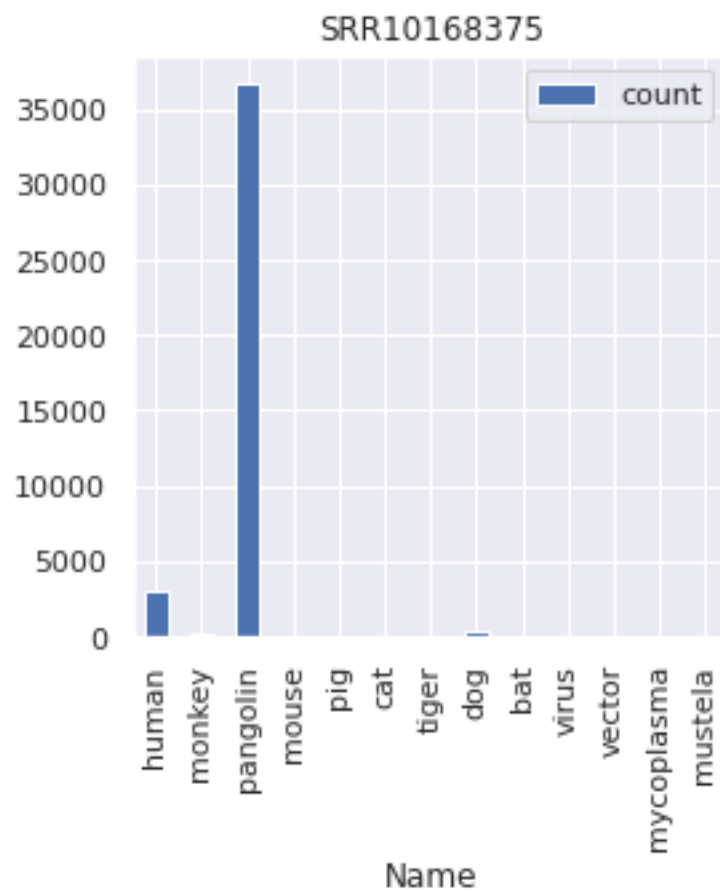
```

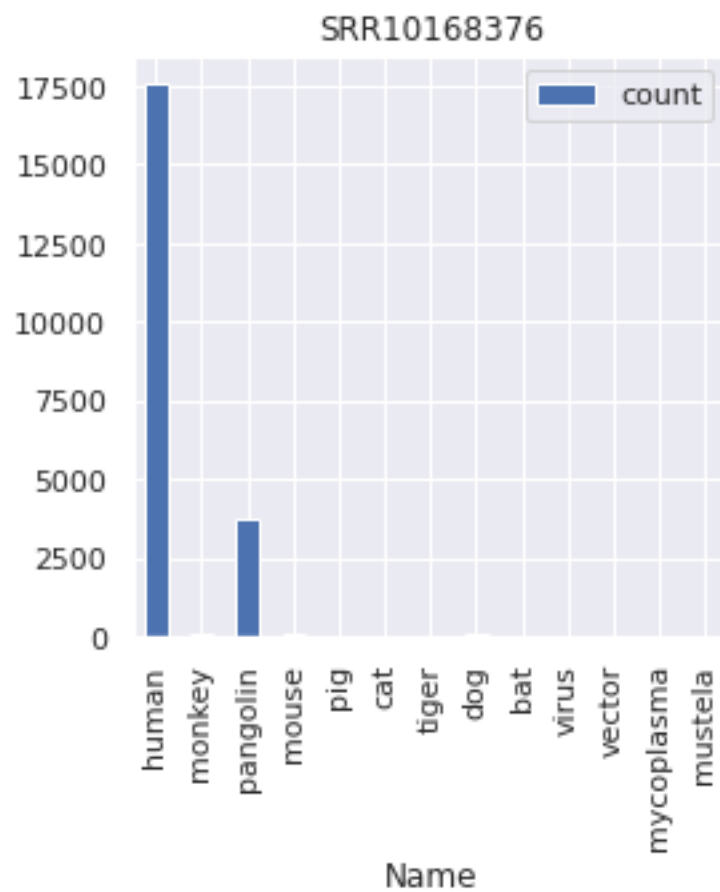
[14]: for sra in sra_list:
    try:
        df=get_descr(sra)
        df.to_csv(BASE_PATH+sra+'/magic_blast/
↪'+f'{sra}_{dbname}_{kmer}_magicBLAST_species_df.csv')
        plot_df(df, sra)
    except FileNotFoundError:
        pass
    except AttributeError:
        pass

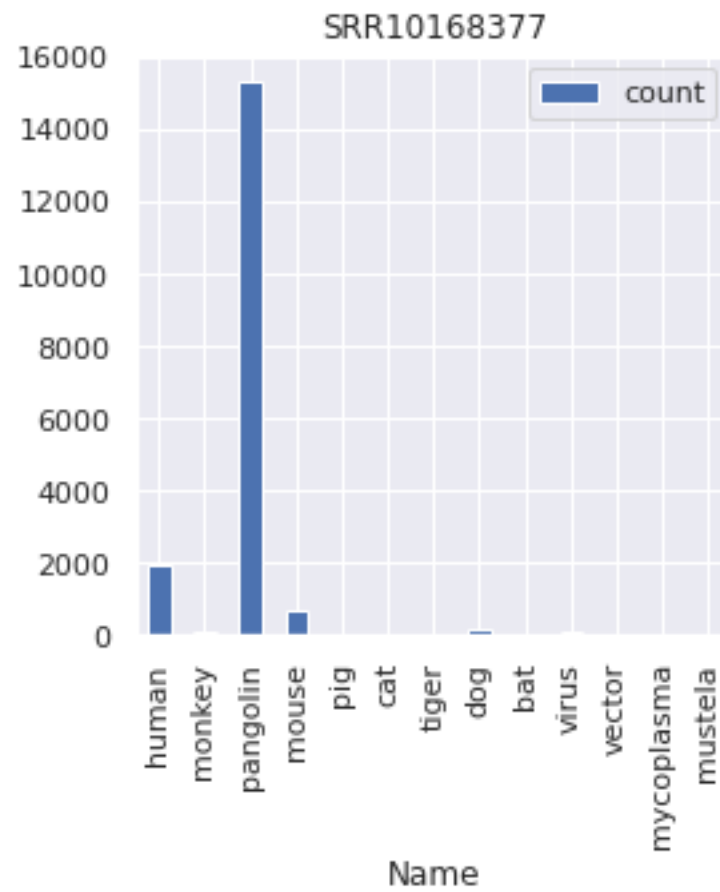
```

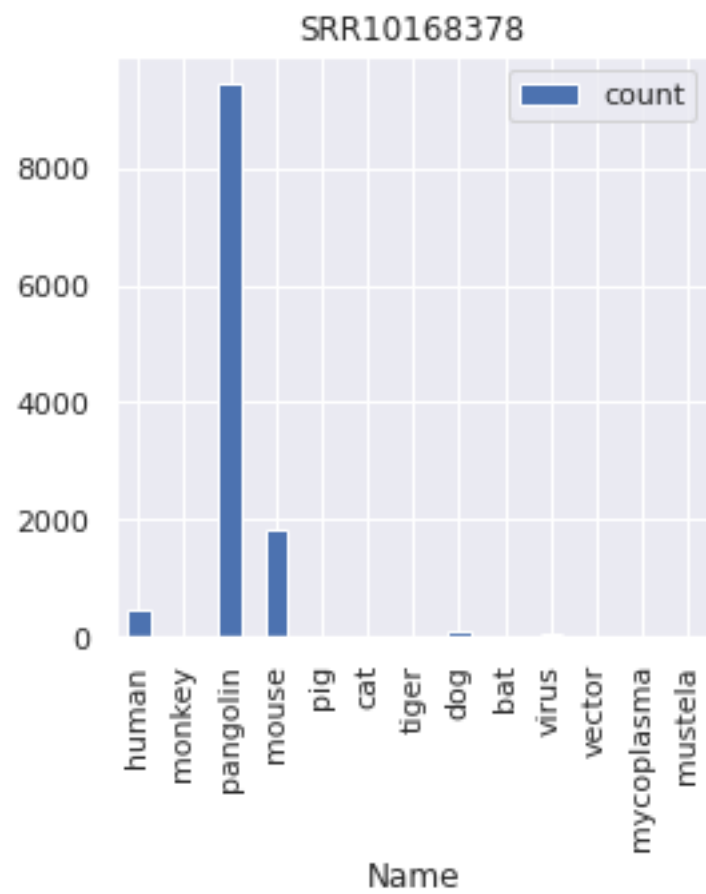


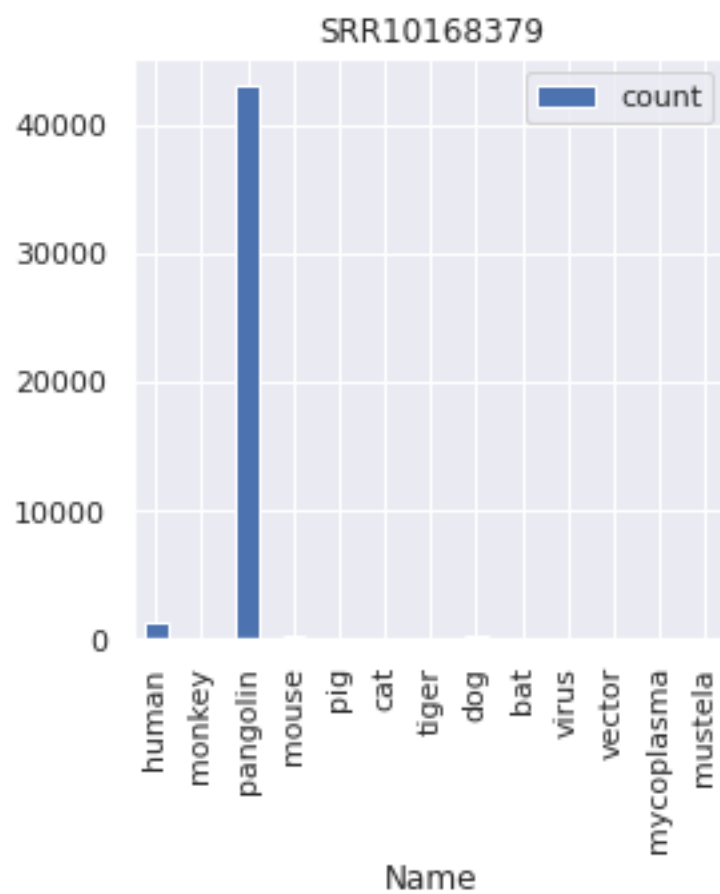


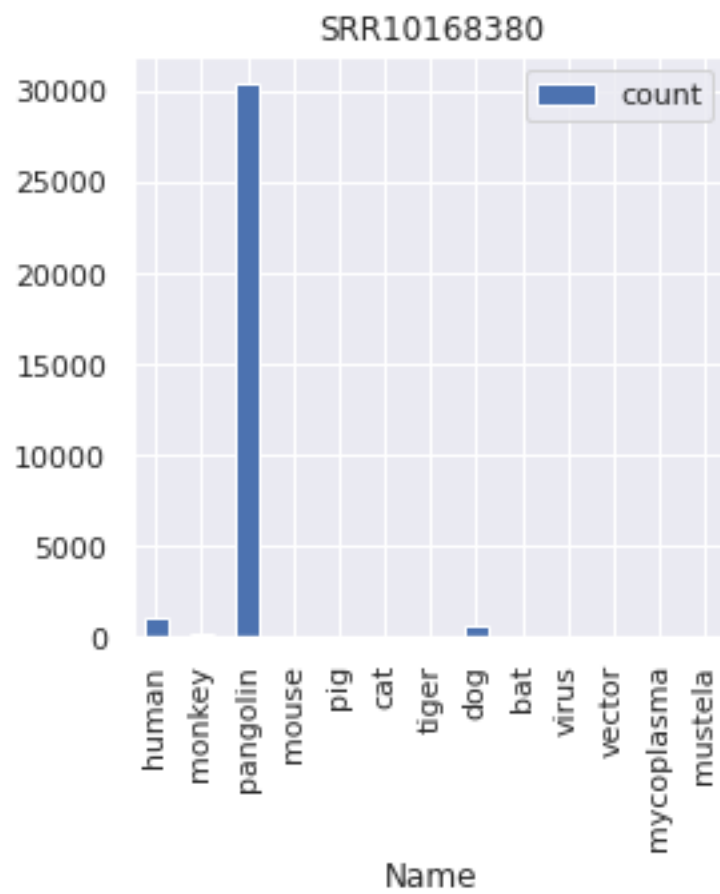


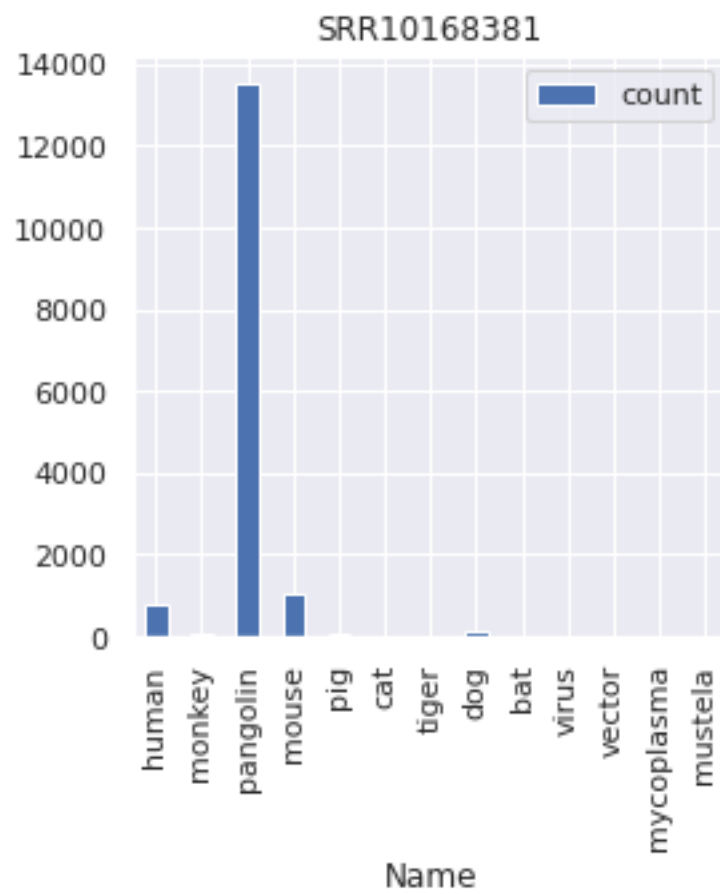


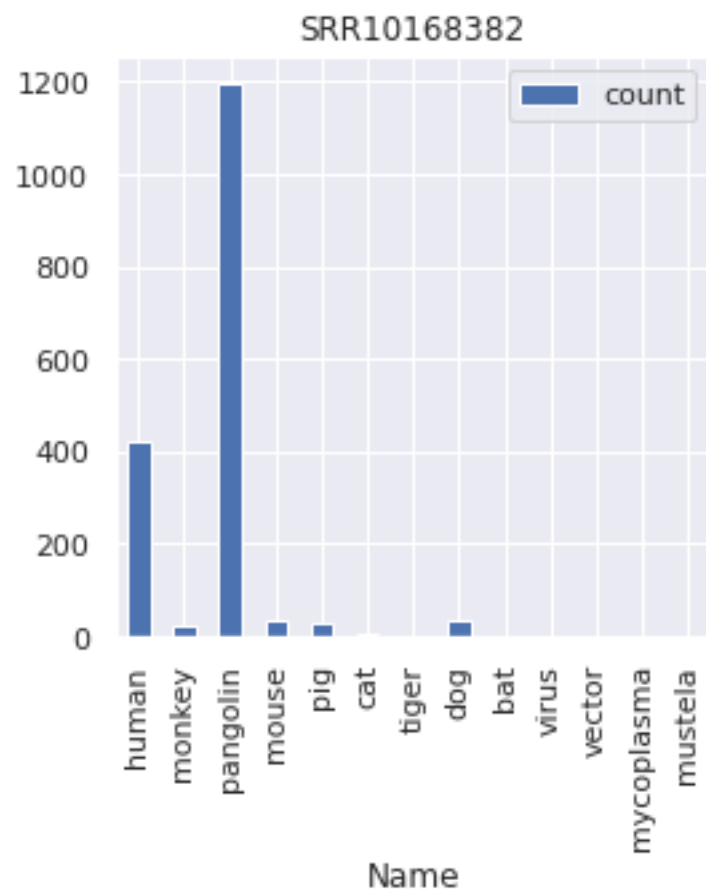


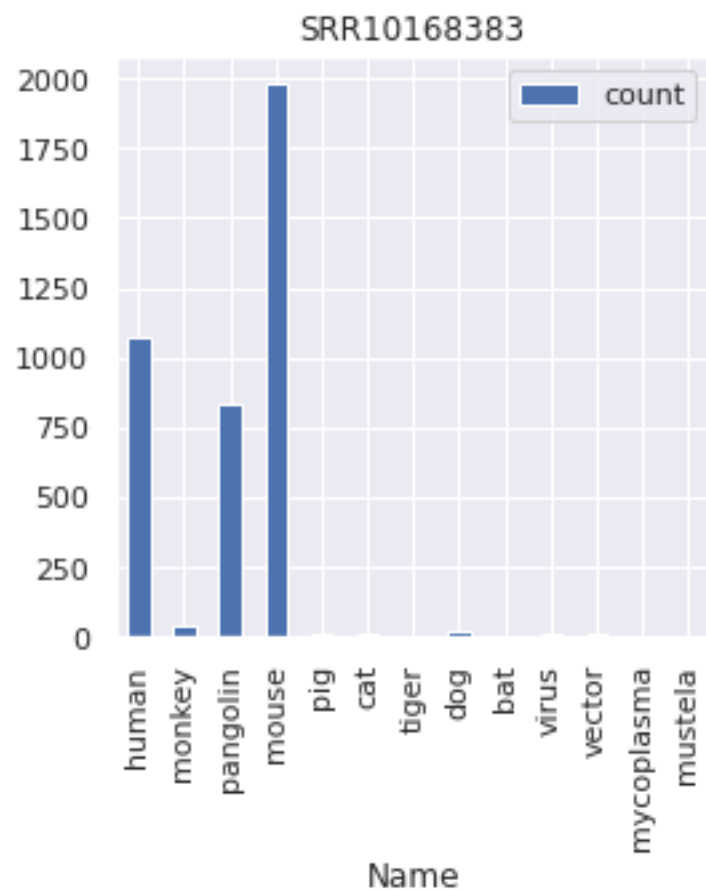


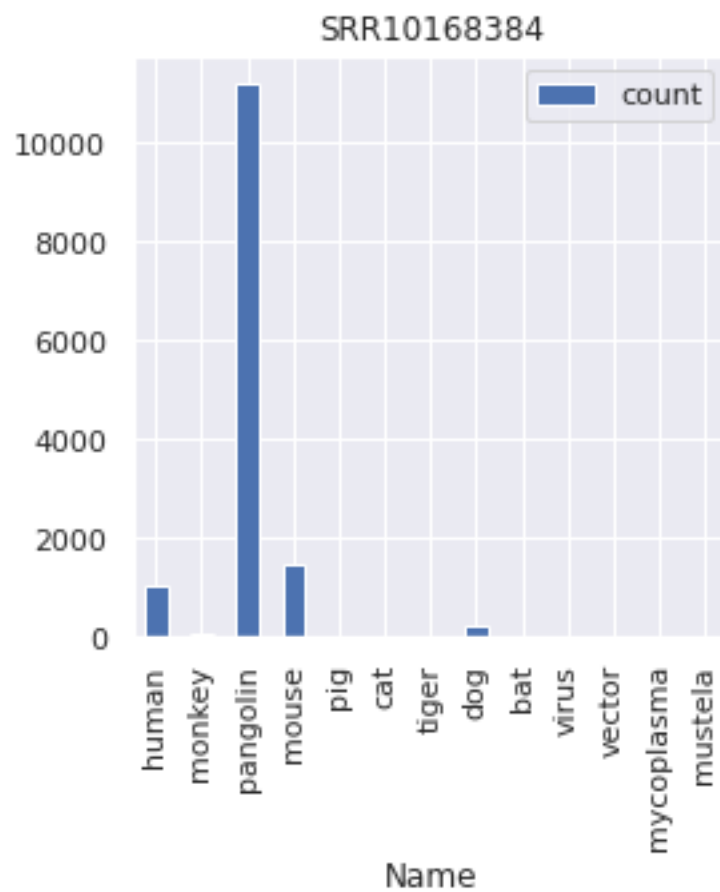


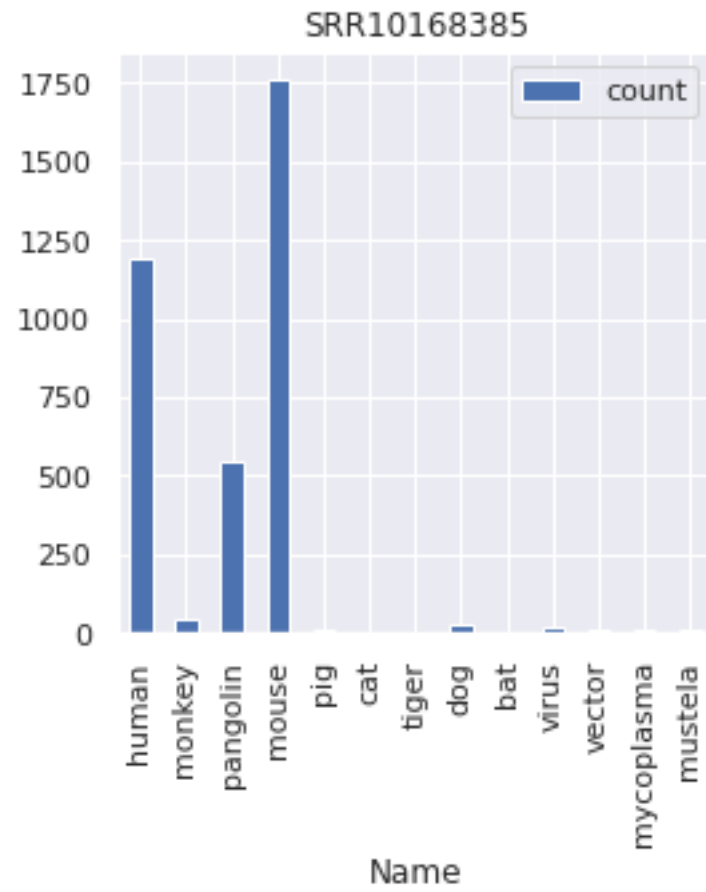


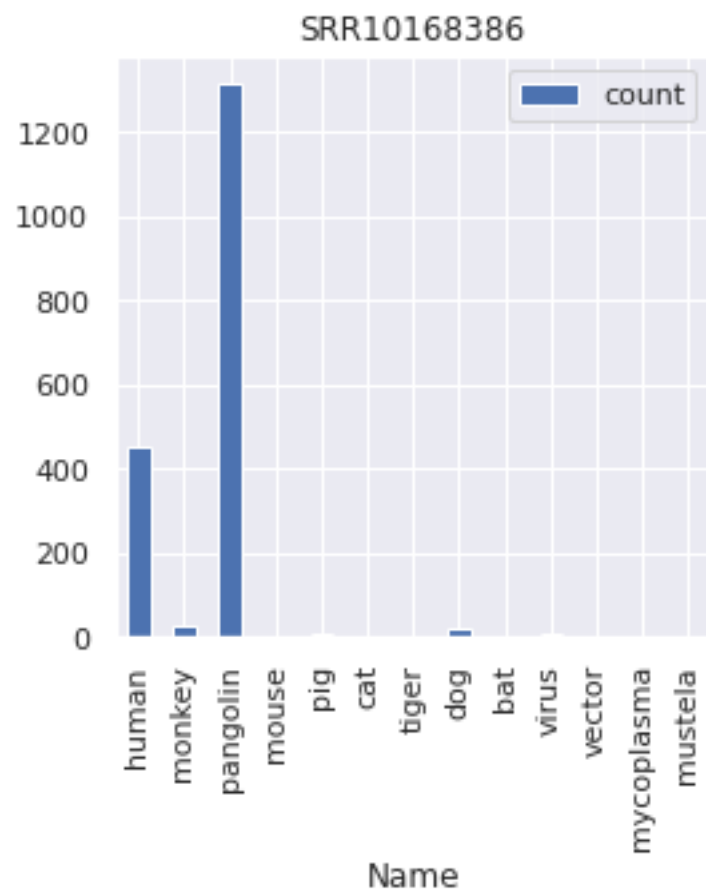


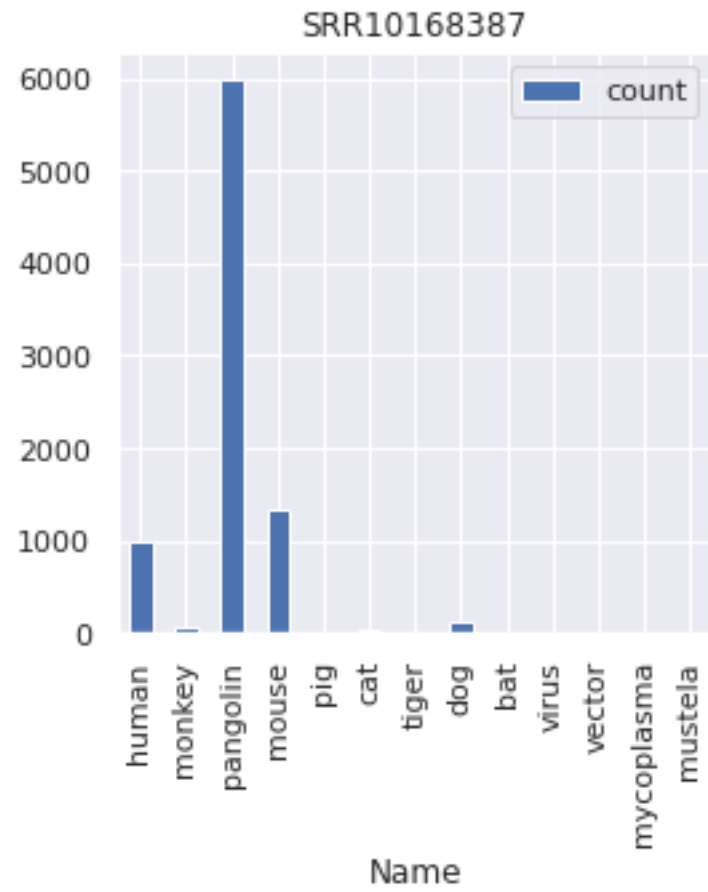


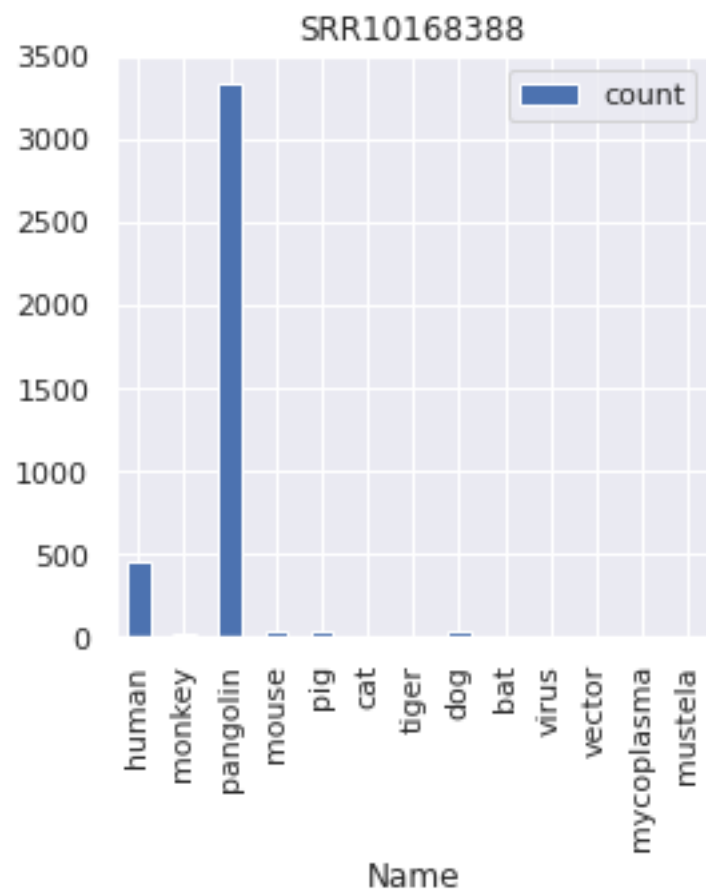


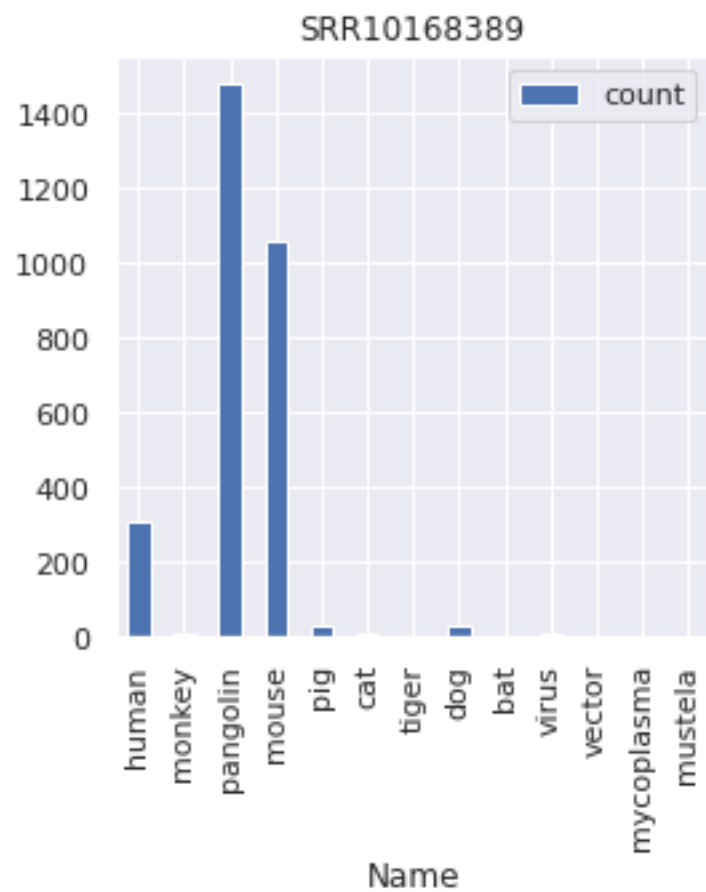


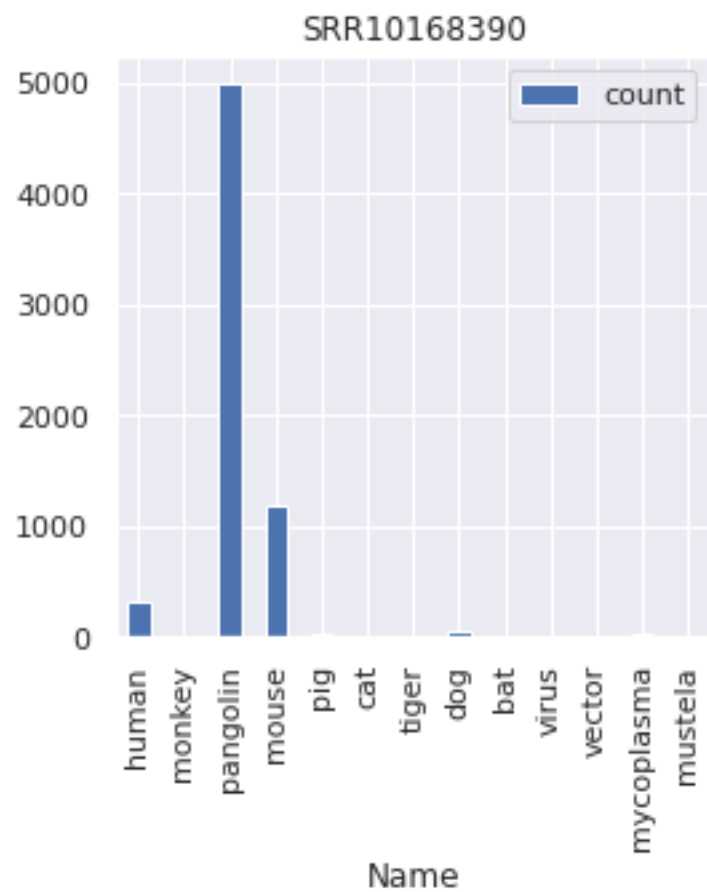


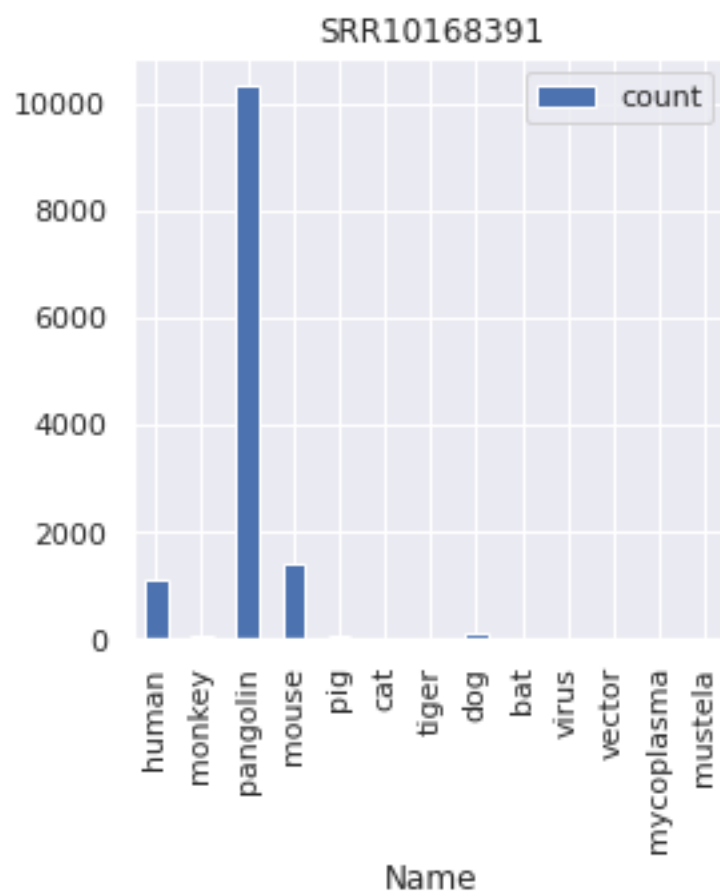


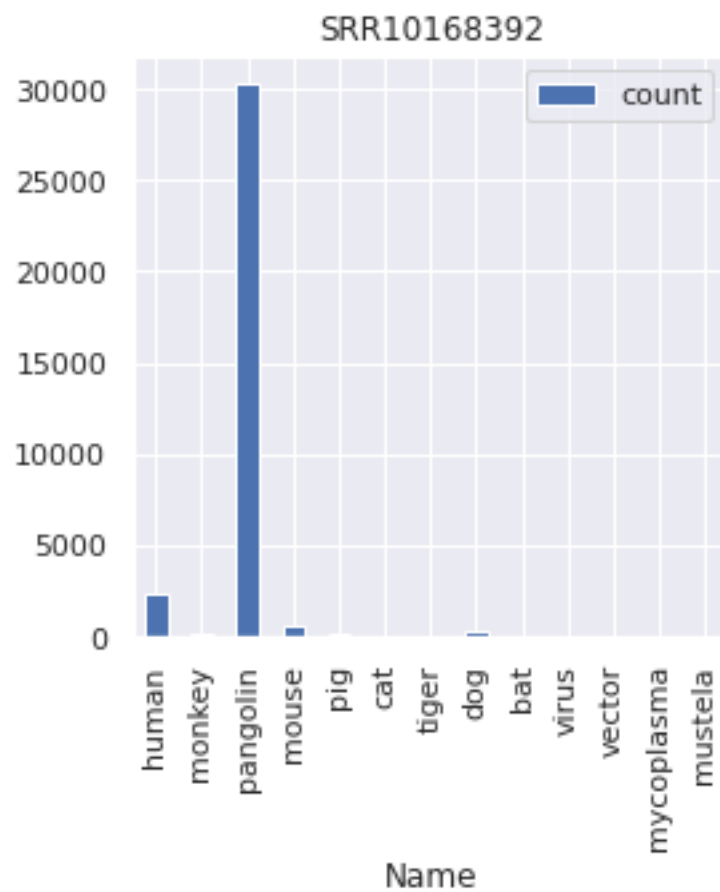


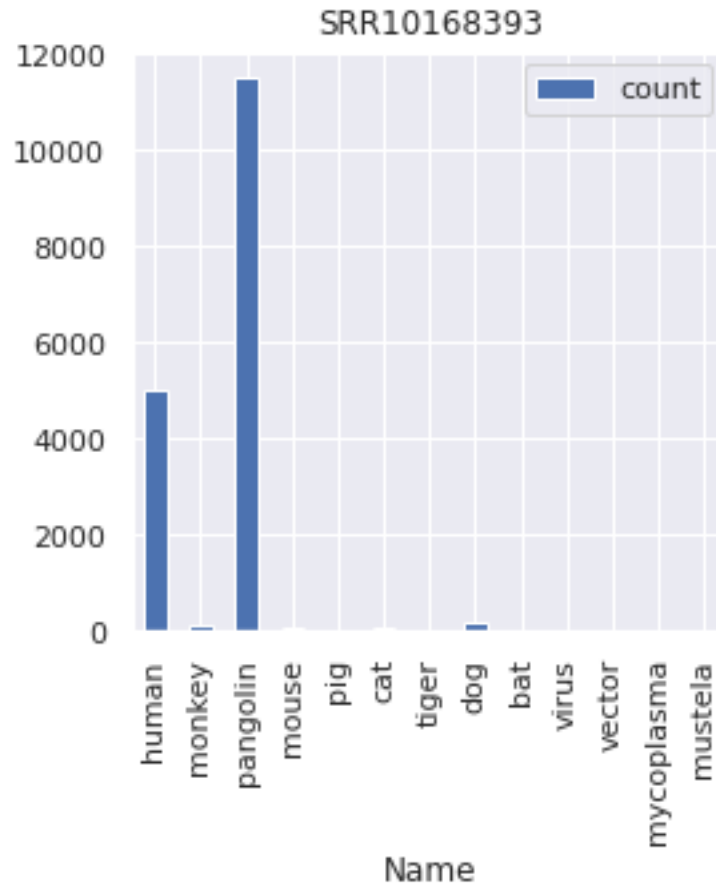












```
[15]: pathlib.Path(BASE_PATH+'general_plots/').mkdir(exist_ok=True)
```

```
def multi_plot():
    fig, axis = plt.subplots(2, 5, figsize=(16,8))
    fig.suptitle('Contig counts')
    sra=sra_list[0]
    df=get_descr(sra)
    axis[0,0].bar(df['Name'],df['count'])
    axis[0,0].tick_params(axis='x', rotation=90)
    axis[0,0].set_ylabel('count')
    axis[0,0].title.set_text(sra)

    sra=sra_list[1]
    df=get_descr(sra)
    axis[0,1].bar(df['Name'],df['count'])
    axis[0,1].set_ylabel('count')
    axis[0,1].tick_params(axis='x', rotation=90)
    axis[0,1].set_ylabel('count')
    axis[0,1].title.set_text(sra)
```

```

sra=sra_list[2]
df=get_descr(sra)
axis[0,2].bar(df['Name'],df['count'])
axis[0,2].set_ylabel('count')
axis[0,2].tick_params(axis='x', rotation=90)
axis[0,2].set_ylabel('count')
axis[0,2].title.set_text(sra)

sra=sra_list[3]
df=get_descr(sra)
axis[0,3].bar(df['Name'],df['count'])
axis[0,3].set_ylabel('count')
axis[0,3].tick_params(axis='x', rotation=90)
axis[0,3].set_ylabel('count')
axis[0,3].title.set_text(sra)

sra=sra_list[4]
df=get_descr(sra)
axis[0,4].bar(df['Name'],df['count'])
axis[0,4].set_ylabel('count')
axis[0,4].tick_params(axis='x', rotation=90)
axis[0,4].set_ylabel('count')
axis[0,4].title.set_text(sra)

sra=sra_list[5]
df=get_descr(sra)
axis[1,0].bar(df['Name'],df['count'])
axis[1,0].set_ylabel('count')
axis[1,0].tick_params(axis='x', rotation=90)
axis[1,0].set_ylabel('count')
axis[1,0].title.set_text(sra)

sra=sra_list[6]
df=get_descr(sra)
axis[1,1].bar(df['Name'],df['count'])
axis[1,1].set_ylabel('count')
axis[1,1].tick_params(axis='x', rotation=90)
axis[1,1].set_ylabel('count')
axis[1,1].title.set_text(sra)

sra=sra_list[7]
df=get_descr(sra)
axis[1,2].bar(df['Name'],df['count'])
axis[1,2].set_ylabel('count')
axis[1,2].tick_params(axis='x', rotation=90)

```

```

axis[1,2].set_ylabel('count')
axis[1,2].title.set_text(sra)

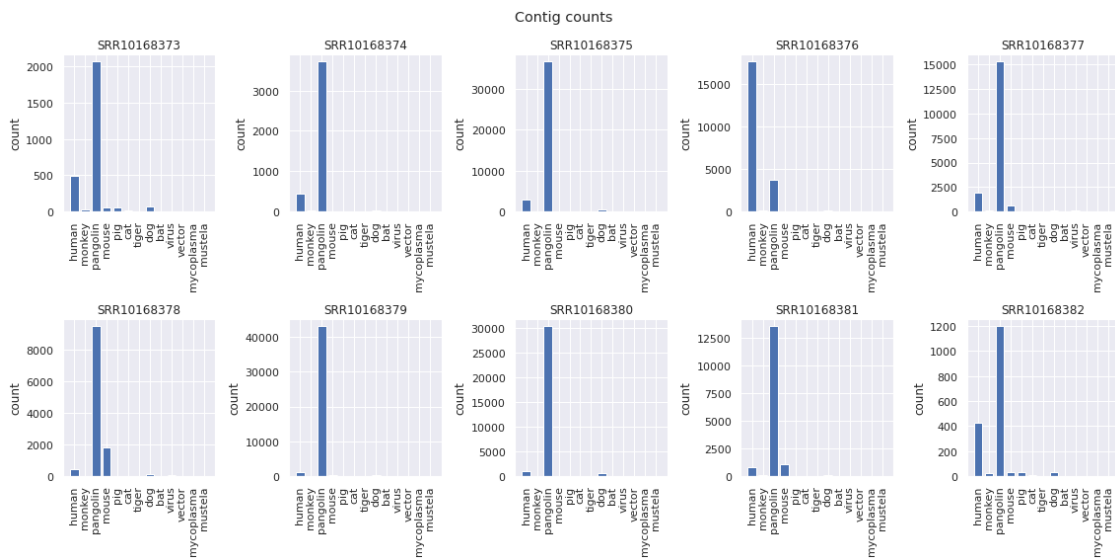
sra=sra_list[8]
df=get_descr(sra)
axis[1,3].bar(df['Name'],df['count'])
axis[1,3].set_ylabel('count')
axis[1,3].tick_params(axis='x', rotation=90)
axis[1,3].set_ylabel('count')
axis[1,3].title.set_text(sra)

sra=sra_list[9]
df=get_descr(sra)
axis[1,4].bar(df['Name'],df['count'])
axis[1,4].set_ylabel('count')
axis[1,4].tick_params(axis='x', rotation=90)
axis[1,4].set_ylabel('count')
axis[1,4].title.set_text(sra)

fig.tight_layout()
fig.savefig(BASE_PATH+'general_plots/
↳'+f'{sra}_{dbname}_{kmer}_magicBLAST_contig_count_matrix.png',
↳bbox_inches="tight")
plt.show()

```

[16]: multi_plot()



```
[17]: frames=[]
      for i,sra in enumerate(sra_list):
          try:
              df = pd.read_csv(BASE_PATH+sra+'/magic_blast/
→'+f'{sra}_{dbname}_{kmer}_magicBLAST_species_df.csv')
              x = df.Name.astype('category')
              df['species_uid'] =x.cat.codes
              df['SRA_val'] =df.SRA.str.strip('SRR')
              df['SRA_val'] = pd.to_numeric(df['SRA_val'])
              frames.append(df)
          except FileNotFoundError:
              pass
      df_sra = pd.concat(frames)
```

```
[18]: df_sra.drop(columns=['Unnamed: 0'],inplace=True)
```

```
[19]: df_sra.head(n=100)
```

```
[19]:
```

	SRA	Name	count	pct_matched	species_uid	SRA_val
0	SRR10168373	human	486	11.18	3	10168373
1	SRR10168373	monkey	21	0.48	4	10168373
2	SRR10168373	pangolin	2067	47.54	8	10168373
3	SRR10168373	mouse	53	1.22	5	10168373
4	SRR10168373	pig	51	1.17	9	10168373
..
4	SRR10168380	pig	52	0.16	9	10168380
5	SRR10168380	cat	66	0.20	1	10168380
6	SRR10168380	tiger	6	0.02	10	10168380
7	SRR10168380	dog	556	1.67	2	10168380
8	SRR10168380	bat	67	0.20	0	10168380

[100 rows x 6 columns]

```
[20]: df_sra.Name.unique()
```

```
[20]: array(['human', 'monkey', 'pangolin', 'mouse', 'pig', 'cat', 'tiger',
        'dog', 'bat', 'virus', 'vector', 'mycoplasma', 'mustela'],
        dtype=object)
```

0.0.3 All nt database matches

```
[21]: total_dict={}
      asc_desc={}
      for sra in sra_list:
          accessions, descriptions, counts, total=get_asc_descr_count(sra)
          for asc,desc,cnt in zip(accessions,descriptions, counts):
              if asc in total_dict:
```

```

        total_dict[asc]+=int(cnt)
    else:
        total_dict[asc]=int(cnt)
    if asc not in asc_desc:
        asc_desc[asc]=desc

```

```
[22]: assert len(total_dict)==len(asc_desc)
```

```
[23]: len(asc_desc)
```

```
[23]: 74327
```

Print out the 100 most common nt database classification for all contigs in the project

```
[24]: def print_top_n_sp(total_dict, max_num):
        listofTuples = sorted(total_dict.items() , reverse=True, key=lambda x: x[1])
        for i, elem in enumerate(listofTuples):
            if i<max_num:
                print(asc_desc[elem[0]], ": " , elem[1] )
            else:
                break

```

```
[25]: print_top_n_sp(total_dict, max_num=10)
```

```

Pseudolabrys sp. FHR47 chromosome, complete genome : 858
Eukaryotic synthetic construct chromosome 16 : 841
Homo sapiens DNA, chromosome 16, nearly complete genome : 810
Lutra lutra genome assembly, chromosome: 16 : 656
Eukaryotic synthetic construct chromosome 17 : 554
Staphylococcus aureus strain WH9628 chromosome : 542
Homo sapiens DNA, chromosome 17, nearly complete genome : 527
Sus scrofa 18S ribosomal RNA gene, complete sequence : 516
Beta vulgaris subsp. vulgaris cultivar KWS2320 chloroplast, complete genome :
507
Homo sapiens clone LA14_101B3 sequence : 506

```

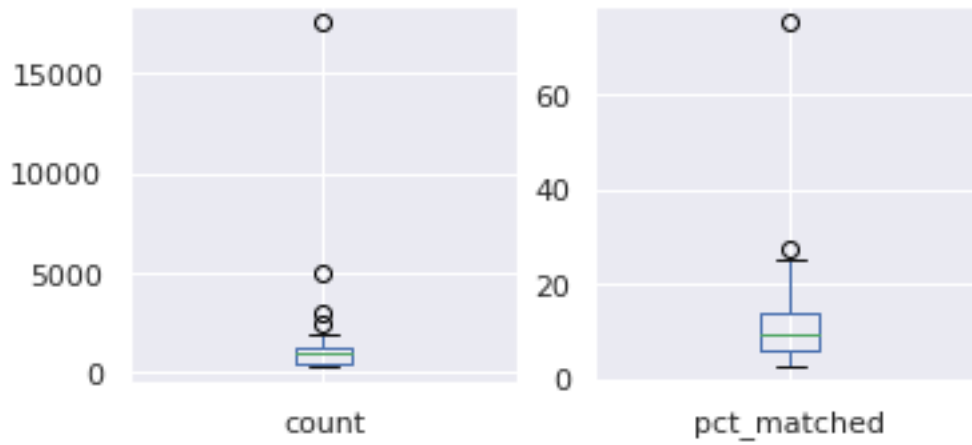
0.0.4 Human

```
[26]: df=df_sra[df_sra['Name'].str.contains('human')]
```

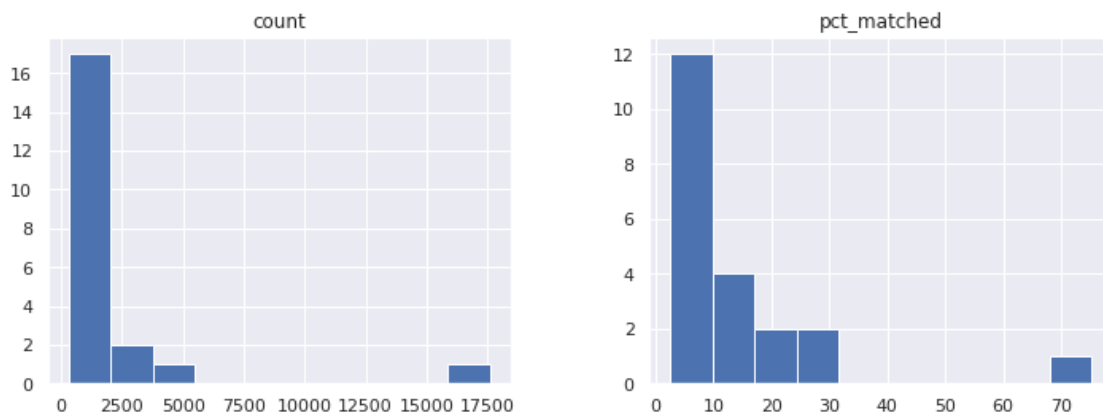
```
[27]: df.drop(columns=['Name'],inplace=True)
sns.set(rc={"figure.figsize":(6, 6)})
df_box=df[['count', 'pct_matched']]
df_box.plot(kind='box', subplots=True, layout=(4,4), sharex=False,
↳sharey=False, figsize=(12,12))

```

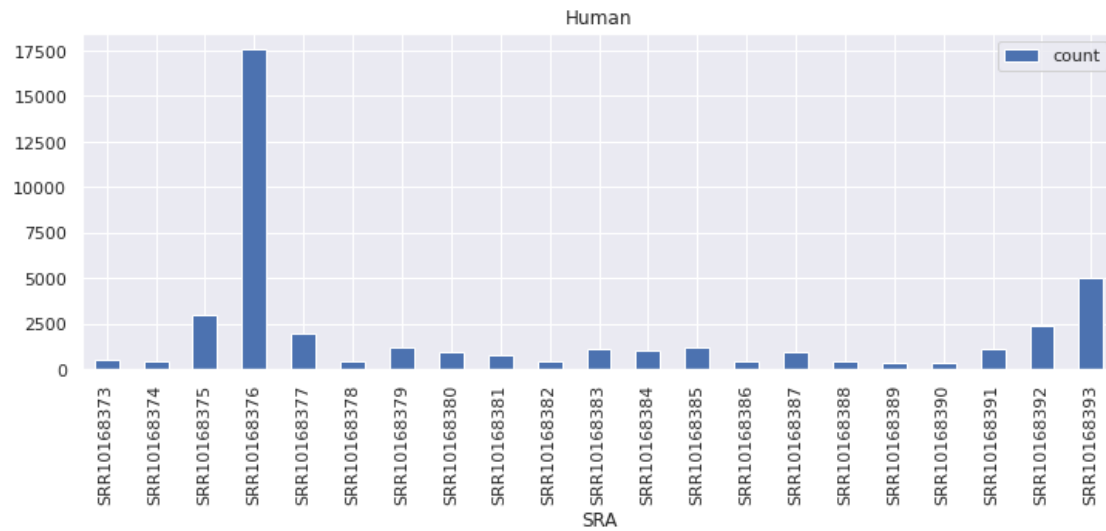
```
[27]: count                AxesSubplot(0.125,0.71587;0.168478x0.16413)
      pct_matched         AxesSubplot(0.327174,0.71587;0.168478x0.16413)
      dtype: object
```



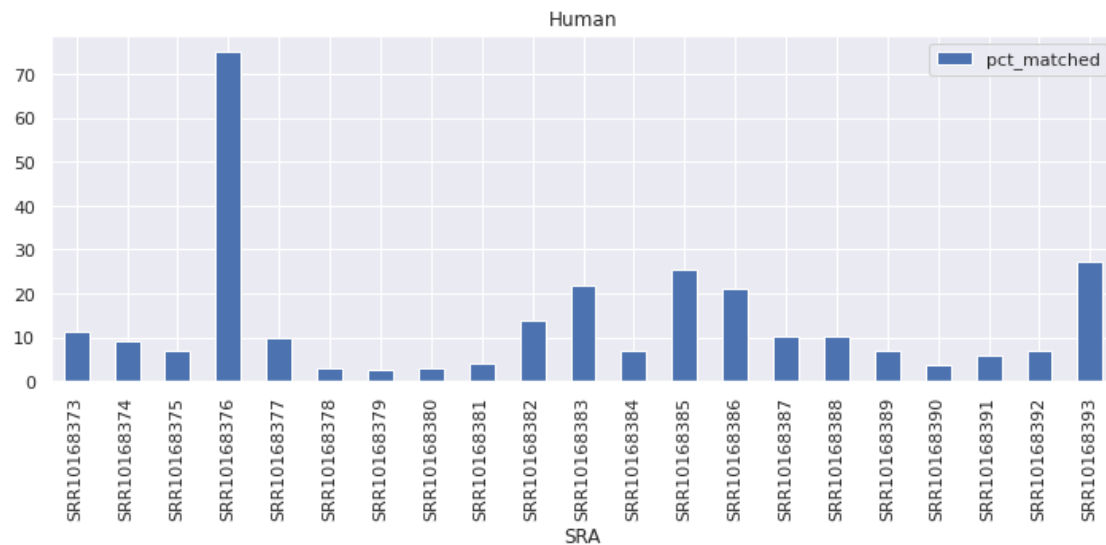
```
[28]: df_box.hist(figsize=(12,4))
      plt.show()
```



```
[29]: ax=df.plot(x='SRA', y='count', kind='bar',figsize=(12,4))
      ax.set_title('Human', fontsize=12)
      plt.show()
```

```
[30]: ax=df.plot(x='SRA', y='pct_matched', kind='bar',figsize=(12,4))
ax.set_title('Human', fontsize=12)
plt.show()
```



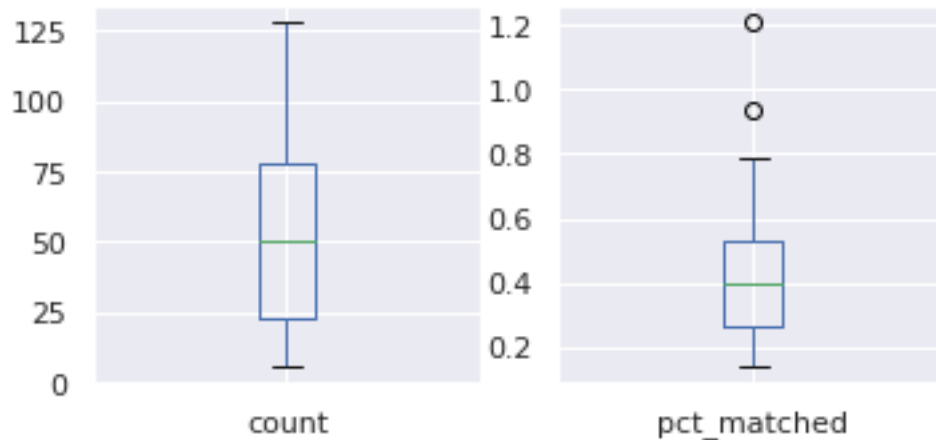
0.0.5 Monkey

```
[31]: s_name='monkey'
```

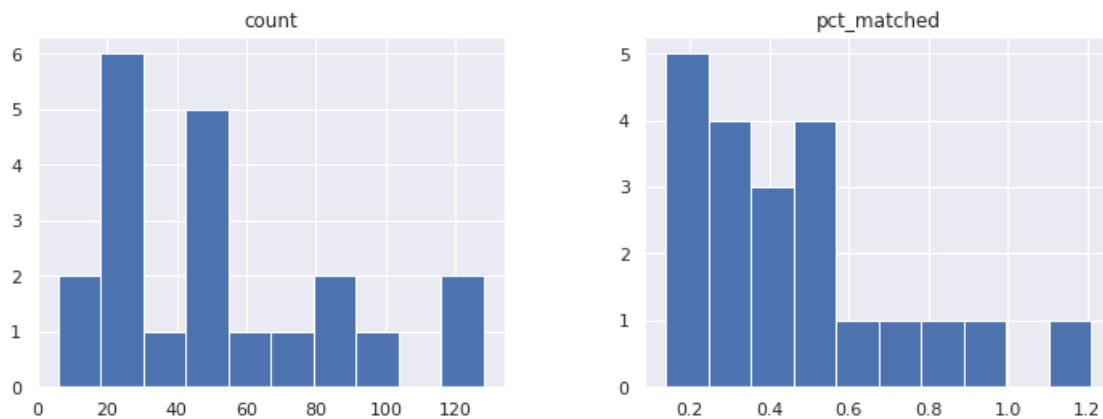
```
[32]: df=df_sra[df_sra['Name'].str.contains(s_name)]
```

```
[33]: df.drop(columns=['Name'],inplace=True)
sns.set(rc={"figure.figsize":(6, 6)})
df_box=df[['count', 'pct_matched']]
df_box.plot(kind='box', subplots=True, layout=(4,4), sharex=False,
→sharey=False, figsize=(12,12))
```

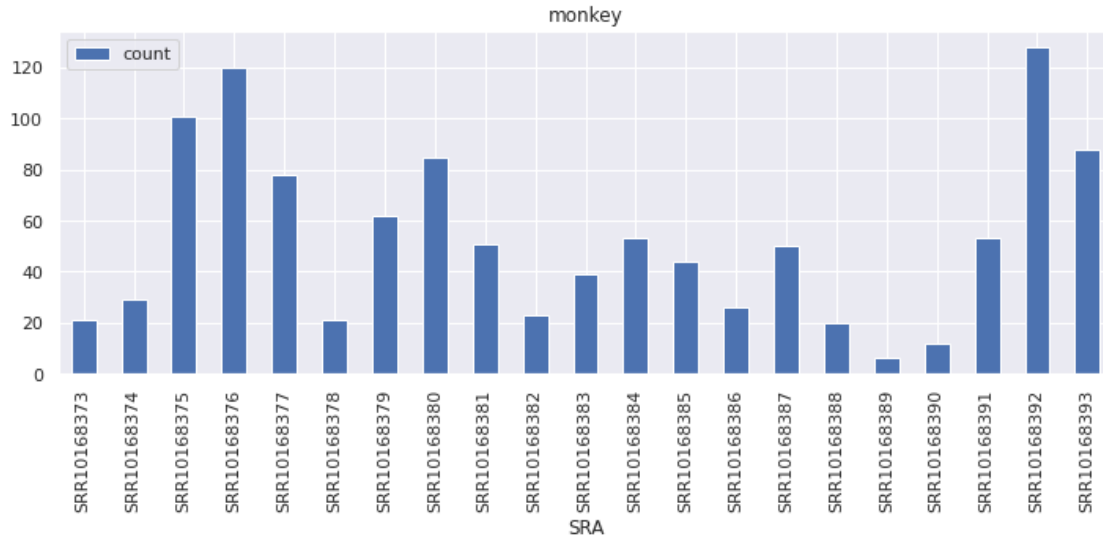
```
[33]: count          AxesSubplot(0.125,0.71587;0.168478x0.16413)
pct_matched      AxesSubplot(0.327174,0.71587;0.168478x0.16413)
dtype: object
```



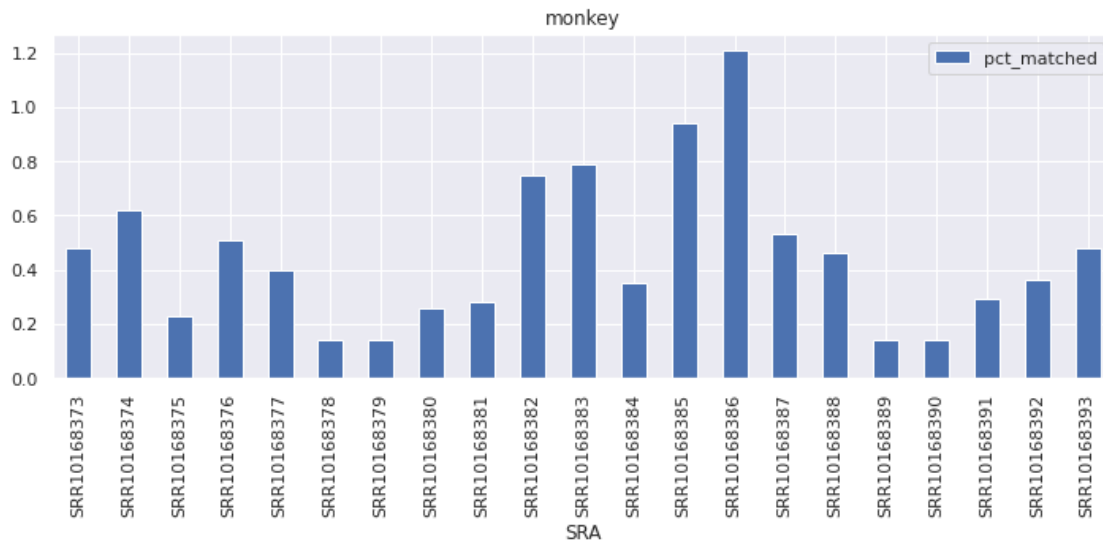
```
[34]: df_box.hist(figsize=(12,4))
plt.show()
```



```
[35]: ax=df.plot(x='SRA', y='count', kind='bar',figsize=(12,4))
ax.set_title(s_name, fontsize=12)
plt.show()
```



```
[36]: ax=df.plot(x='SRA', y='pct_matched', kind='bar',figsize=(12,4))
ax.set_title(s_name, fontsize=12)
plt.show()
```



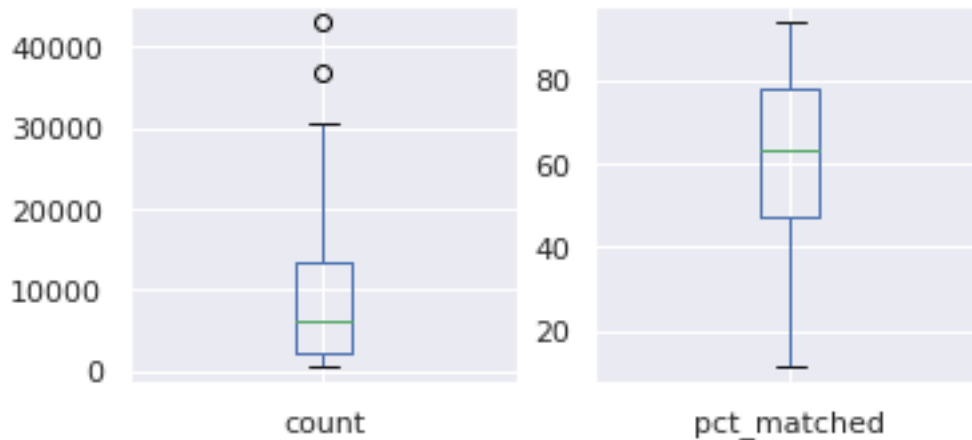
0.0.6 Pangolin

```
[37]: s_name='pangolin'
```

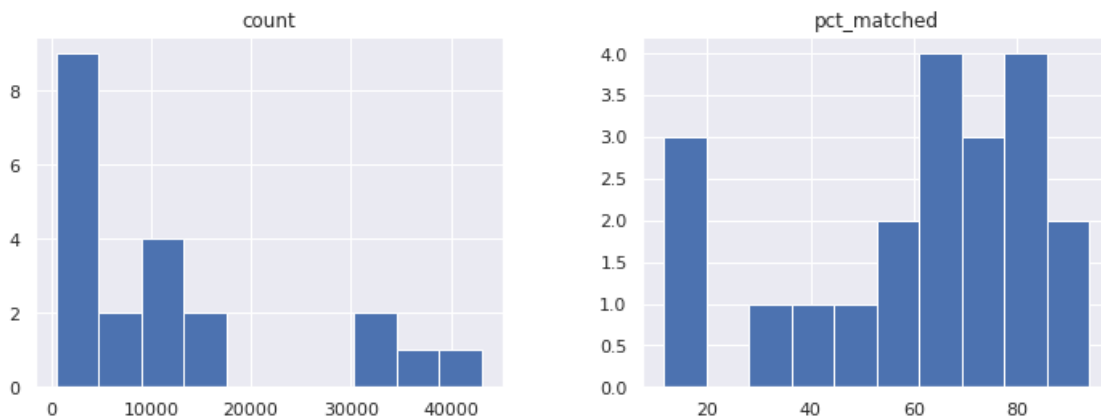
```
[38]: df=df_sra[df_sra['Name'].str.contains(s_name)]
```

```
[39]: df.drop(columns=['Name'],inplace=True)
sns.set(rc={"figure.figsize":(6, 6)})
df_box=df[['count', 'pct_matched']]
df_box.plot(kind='box', subplots=True, layout=(4,4), sharex=False,
→sharey=False, figsize=(12,12))
```

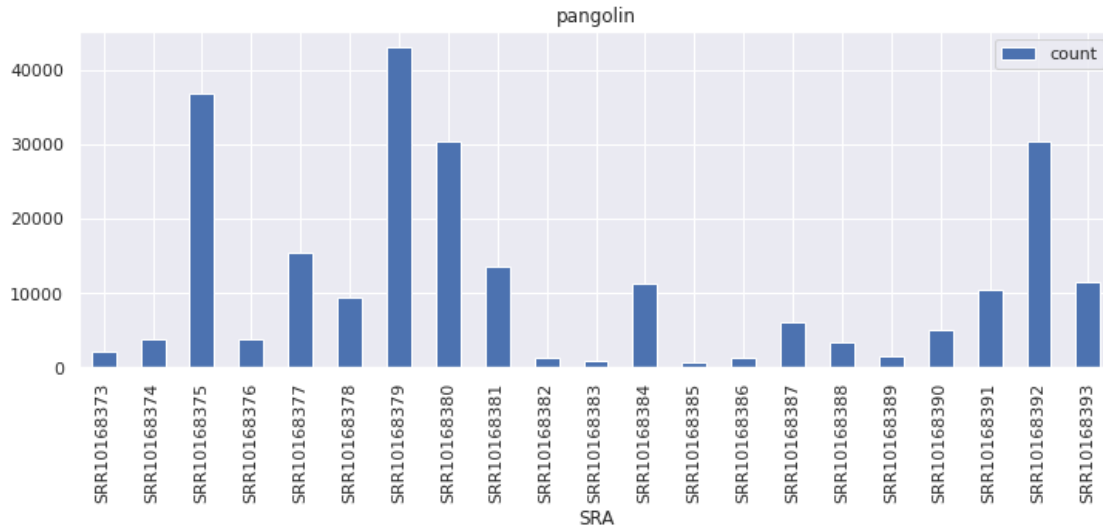
```
[39]: count          AxesSubplot(0.125,0.71587;0.168478x0.16413)
pct_matched      AxesSubplot(0.327174,0.71587;0.168478x0.16413)
dtype: object
```



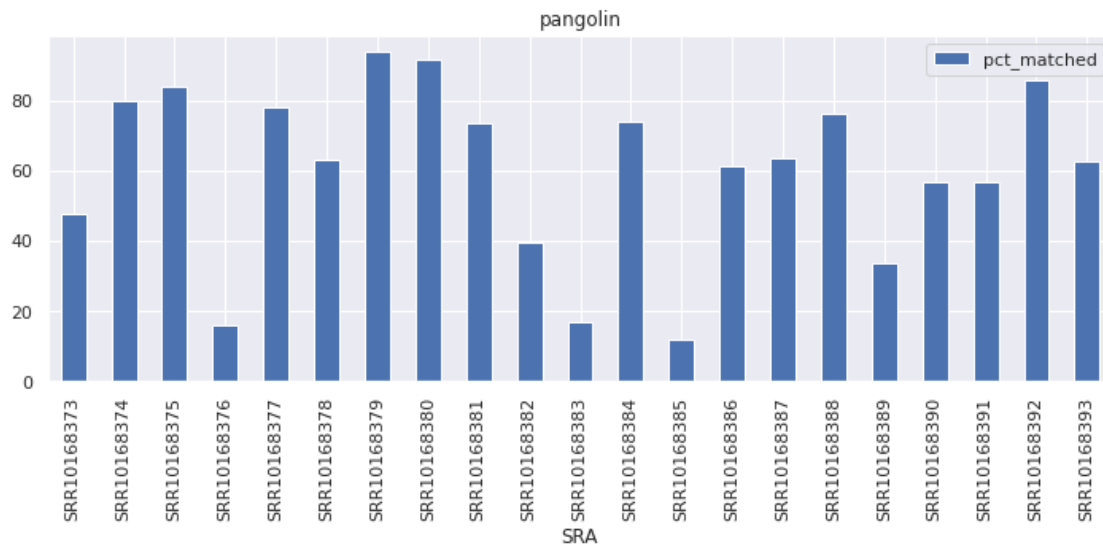
```
[40]: df_box.hist(figsize=(12,4))
plt.show()
```



```
[41]: ax=df.plot(x='SRA', y='count', kind='bar',figsize=(12,4))
ax.set_title(s_name, fontsize=12)
plt.show()
```



```
[42]: ax=df.plot(x='SRA', y='pct_matched', kind='bar',figsize=(12,4))
ax.set_title(s_name, fontsize=12)
plt.show()
```



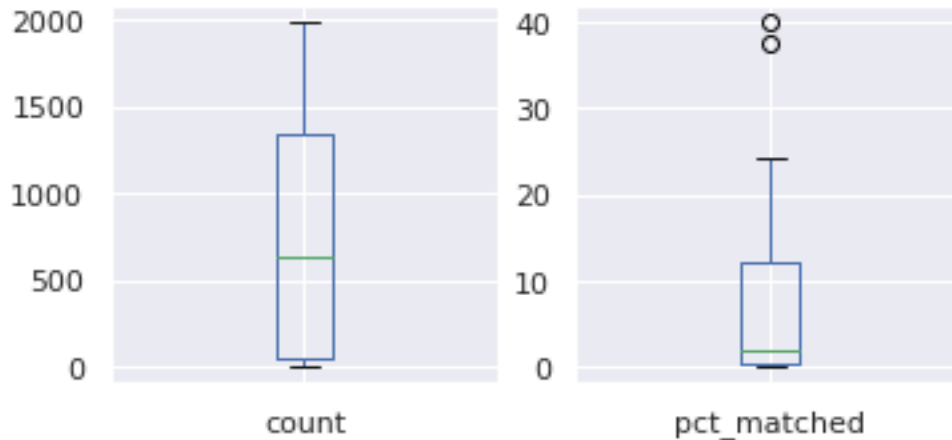
0.0.7 Mouse

```
[43]: s_name='mouse'
```

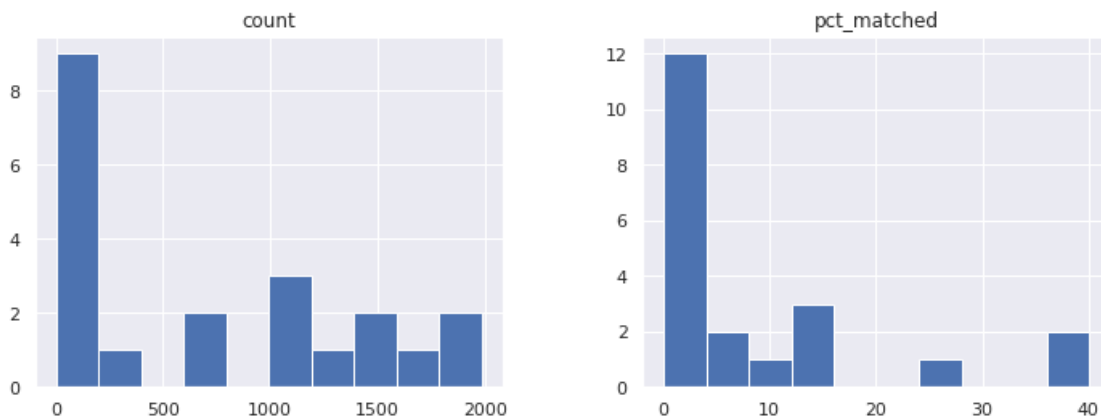
```
[44]: df=df_sra[df_sra['Name'].str.contains(s_name)]
```

```
[45]: df.drop(columns=['Name'],inplace=True)
sns.set(rc={"figure.figsize":(6, 6)})
df_box=df[['count', 'pct_matched']]
df_box.plot(kind='box', subplots=True, layout=(4,4), sharex=False,
→sharey=False, figsize=(12,12))
```

```
[45]: count      AxesSubplot(0.125,0.71587;0.168478x0.16413)
pct_matched    AxesSubplot(0.327174,0.71587;0.168478x0.16413)
dtype: object
```



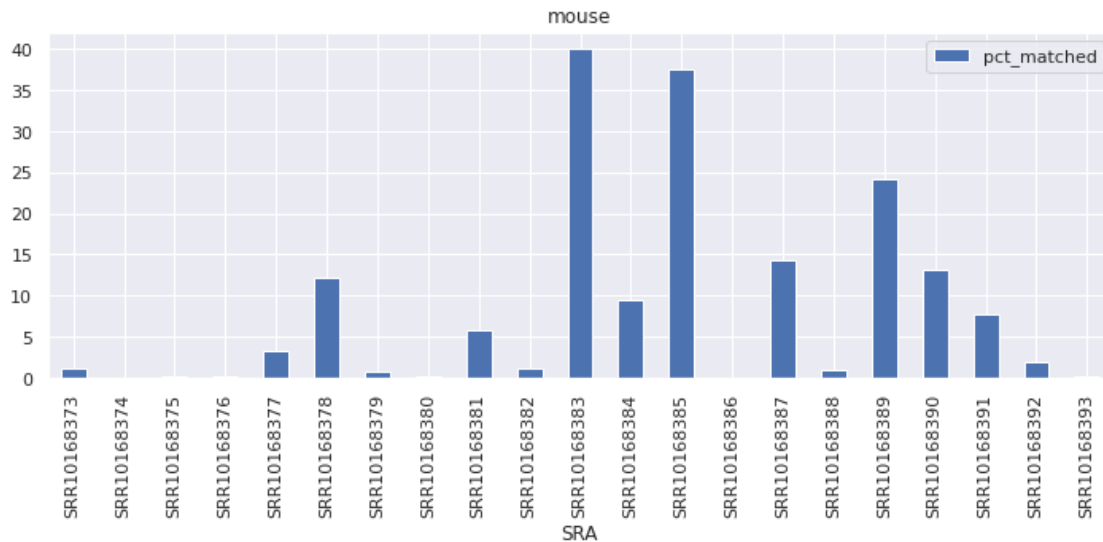
```
[46]: df_box.hist(figsize=(12,4))
plt.show()
```



```
[47]: ax=df.plot(x='SRA', y='count', kind='bar',figsize=(12,4))
ax.set_title(s_name, fontsize=12)
plt.show()
```



```
[48]: ax=df.plot(x='SRA', y='pct_matched', kind='bar',figsize=(12,4))
ax.set_title(s_name, fontsize=12)
plt.show()
```



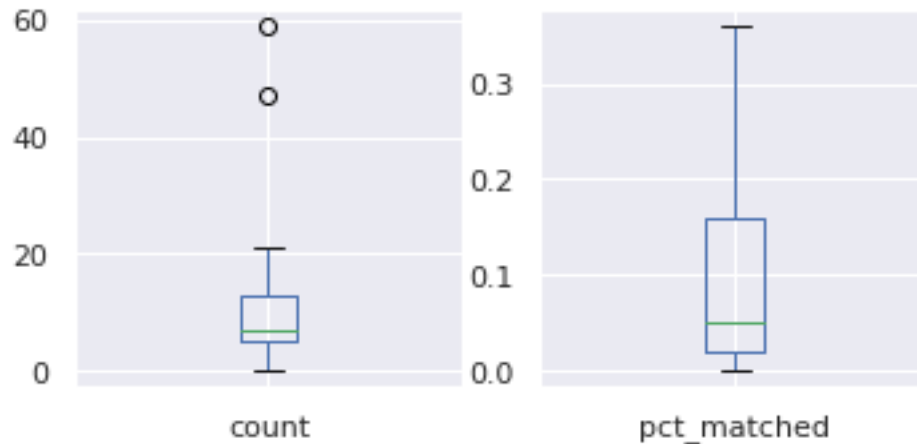
0.0.8 virus

```
[49]: s_name='virus'
```

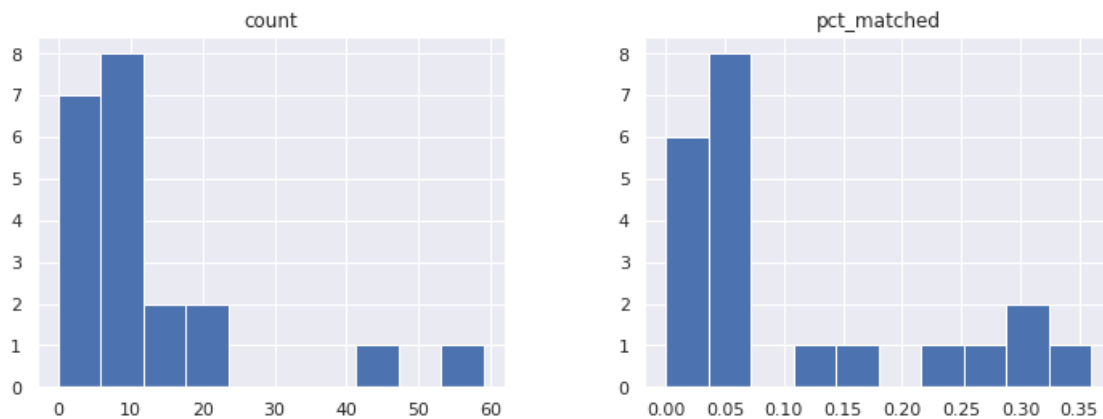
```
[50]: df=df_sra[df_sra['Name'].str.contains(s_name)]
```

```
[51]: df.drop(columns=['Name'],inplace=True)
sns.set(rc={"figure.figsize":(6, 6)})
df_box=df[['count', 'pct_matched']]
df_box.plot(kind='box', subplots=True, layout=(4,4), sharex=False,
→sharey=False, figsize=(12,12))
```

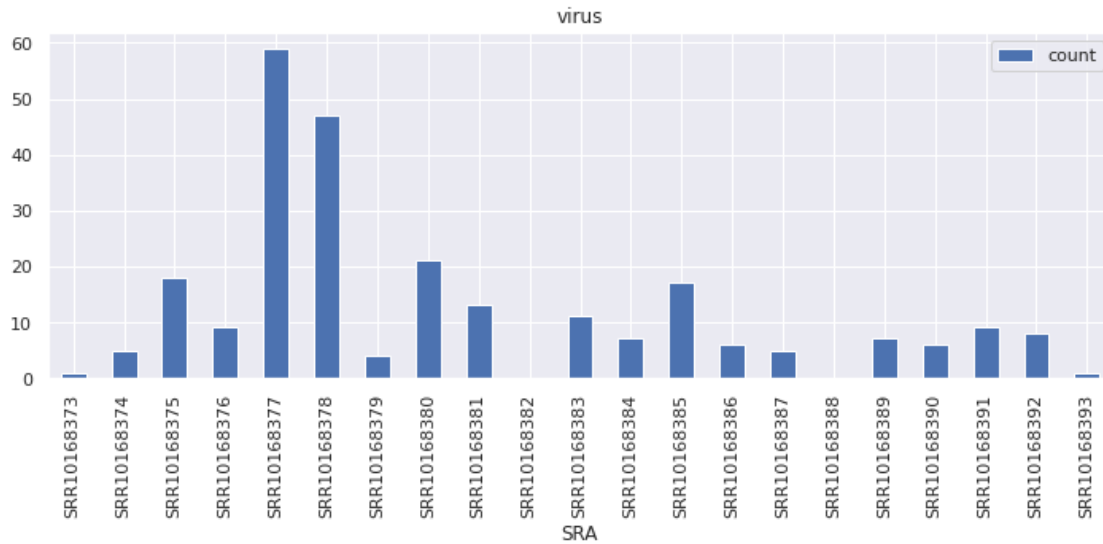
```
[51]: count          AxesSubplot(0.125,0.71587;0.168478x0.16413)
pct_matched        AxesSubplot(0.327174,0.71587;0.168478x0.16413)
dtype: object
```



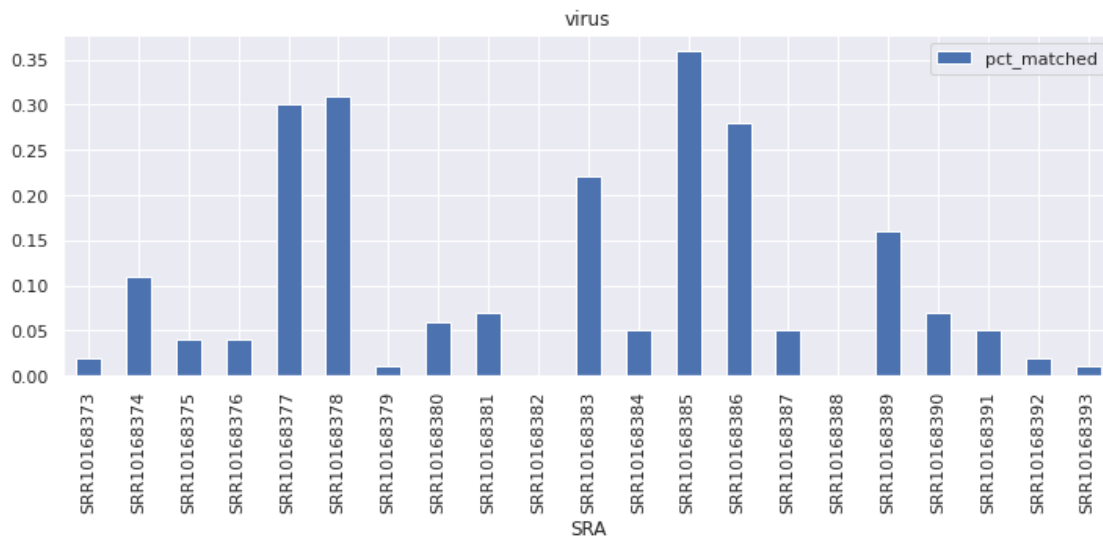
```
[52]: df_box.hist(figsize=(12,4))
plt.show()
```



```
[53]: ax=df.plot(x='SRA', y='count', kind='bar',figsize=(12,4))
ax.set_title(s_name, fontsize=12)
plt.show()
```

```
[54]: ax=df.plot(x='SRA', y='pct_matched', kind='bar',figsize=(12,4))
ax.set_title(s_name, fontsize=12)
plt.show()
```



0.0.9 Top nt database contigs matches per SRA

```
[55]: def get_sra_dict(sra):
total_dict={}
asc_desc={}
accessions, descriptions, counts, total=get_asc_descr_count(sra)
```

```

for asc,desc,cnt in zip(accessions,descriptions, counts):
    if asc in total_dict:
        total_dict[asc]+=int(cnt)
    else:
        total_dict[asc]=int(cnt)
    if asc not in asc_desc:
        asc_desc[asc]=desc
return total_dict, asc_desc, total

```

```

[56]: for sra in sra_list:
    total_dict, asc_desc, total= get_sra_dict(sra)
    print(f'{sra}, number of contigs {total}')
    print_top_n_sp(total_dict, max_num=10)
    print('\n')

```

```

SRR10168373, number of contigs 2117
Manis javanica isolate EP2 mitochondrion, complete genome : 151
Manis javanica isolate T298 mitochondrion, complete genome : 150
Manis javanica isolate MJA633 mitochondrion, complete genome : 140
Manis pentadactyla mitochondrion, complete genome : 101
Manis javanica isolate MP_PG03-UM mitochondrion, complete genome : 100
Sphingomonas paucimobilis strain FDAARGOS_908 chromosome, complete genome : 62
Sphingomonas paucimobilis strain FDAARGOS_881 chromosome, complete genome : 58
Lutra lutra genome assembly, chromosome: 16 : 50
Sphingomonas sp. LK11, complete genome : 49
Staphylococcus aureus strain WH9628 chromosome : 49

```

```

SRR10168374, number of contigs 3784
Fusobacterium varium ATCC 27725 chromosome, complete genome : 10
Fusobacterium varium strain NCTC10560 genome assembly, chromosome: 1 : 10
Fusobacterium ulcerans ATCC 49185 isolate Fusobacterium ulcerans 81A6 genome
assembly, chromosome: 1 : 10
PREDICTED: Manis javanica AHNAK nucleoprotein (AHNAK), mRNA : 10
Fusobacterium ulcerans strain NCTC12112 genome assembly, chromosome: 1 : 9
Peptostreptococcus anaerobius ATCC:27337 16S-23S ribosomal intergenic spacer and
23S ribosomal RNA gene, partial sequence : 8
Fusobacterium varium Fv113-g1 DNA, complete genome : 7
Acinetobacter towneri strain 19110F47 chromosome, complete genome : 7
PREDICTED: Manis javanica thrombospondin 1 (THBS1), mRNA : 7
Staphylococcus condimentii strain DSM 11674, complete genome : 6

```

```

SRR10168375, number of contigs 24599
Homo sapiens DNA, chromosome 16, nearly complete genome : 153
Eukaryotic synthetic construct chromosome 16 : 151
Homo sapiens interleukin 9 receptor (IL9Rps) pseudogene, complete sequence; and

```

DNA-directed RNA polymerases III 12.5 kDa polypeptide (POLR3K), U11/U12 snRNP 25 kDa protein (C16orf33), RHBDG1 (C16orf8), DNA-3-methyladenine glycosylase (MPG), -14 gene protein (C16orf35), hemoglobin zeta subunit (HBZ), HBD (HBD), alpha globin (HBA2), alpha globin (HBA1), theta globin (HBQ1), and putative RNA-binding protein Luc7-like 1 (LUC7L) genes, complete cds : 73
Homo sapiens 16p13.3 sequence section 1 of 8 : 70
Homo sapiens 16p13.3 sequence section 2 of 8 : 60
Propionibacterium acnes TypeIA2 P.acn17, complete genome : 60
Propionibacterium acnes TypeIA2 P.acn31, complete genome : 58
Cutibacterium acnes TP-CU389 DNA, complete genome : 56
Propionibacterium acnes TypeIA2 P.acn33, complete genome : 56
Lutra lutra genome assembly, chromosome: 16 : 53

SRR10168376, number of contigs 7147

Eukaryotic synthetic construct chromosome 16 : 414
Homo sapiens DNA, chromosome 16, nearly complete genome : 382
Eukaryotic synthetic construct chromosome 19 : 297
Eukaryotic synthetic construct chromosome 17 : 292
Homo sapiens DNA, chromosome 17, nearly complete genome : 289
Homo sapiens DNA, chromosome 19, nearly complete genome : 272
Eukaryotic synthetic construct chromosome 19 : 158
Homo sapiens titin (TTN) gene, complete cds : 115
Homo sapiens DNA, chromosome 20, nearly complete genome : 114
Homo sapiens DNA, chromosome 15, nearly complete genome : 113

SRR10168377, number of contigs 13755

Psychrobacter sp. PRwf-1 chromosome, complete genome : 21
Eukaryotic synthetic construct chromosome 17 : 21
Lutra lutra genome assembly, chromosome: 16 : 21
Homo sapiens DNA, chromosome 17, nearly complete genome : 21
Pangolin coronavirus isolate MP789, complete genome : 20
Staphylococcus aureus strain WH9628 chromosome : 18
PREDICTED: Manis javanica AHNAK nucleoprotein (AHNAK), mRNA : 18
Sus scrofa 18S ribosomal RNA gene, complete sequence : 18
Homo sapiens clone LA14_101B3 sequence : 18
Mouse DNA sequence from clone RP23-81C12 on chromosome 17, complete sequence : 18

SRR10168378, number of contigs 11263

Sporosarcina ureilytica strain LMG 22257 chromosome, complete genome : 55
Sporosarcina pasteurii strain BNCC 337394 chromosome, complete genome : 35
Staphylococcus arlettae P2 DNA, complete genome : 34
Bacillus sp. OxB-1 DNA, complete genome : 31
Vagococcus carniphilus strain ATCC BAA-640 chromosome, complete genome : 30
Hathewayia histolytica strain NCTC503 genome assembly, chromosome: 1 : 25

Savagea faecisuis strain Tyl34 23S ribosomal RNA gene, partial sequence : 24
Savagea faecisuis strain Tyl40 23S ribosomal RNA gene, partial sequence : 24
Psychrobacter sp. PRwf-1 chromosome, complete genome : 22
Lutra lutra genome assembly, chromosome: 16 : 21

SRR10168379, number of contigs 25159

PREDICTED: Manis javanica AHNAK nucleoprotein (AHNAK), mRNA : 33
Roseomonas mucosa strain AD2 chromosome, complete genome : 27
Lutra lutra genome assembly, chromosome: 16 : 25
Roseomonas mucosa strain FDAARGOS_658 chromosome 4, complete sequence : 23
Roseomonas sp. FDAARGOS_362 chromosome, complete genome : 21
PREDICTED: Manis javanica dystonin (DST), transcript variant X2, mRNA : 20
PREDICTED: Manis javanica igE-binding protein-like (LOC118971992), partial mRNA : 18
PREDICTED: Manis javanica dystonin (DST), transcript variant X3, mRNA : 18
PREDICTED: Manis javanica dystonin (DST), transcript variant X4, mRNA : 18
PREDICTED: Manis javanica dystonin (DST), transcript variant X5, mRNA : 18

SRR10168380, number of contigs 19957

Lutra lutra genome assembly, chromosome: 16 : 49
Homo sapiens clone N29M24 sequence : 37
Homo sapiens clone LA14_101B3 sequence : 37
Homo sapiens clone LA13_165F6 sequence : 37
Mus musculus clone contig 1 chromocenter region genomic sequence : 36
Staphylococcus aureus strain WH9628 chromosome : 36
Homo sapiens clone LA15_25H3 sequence : 36
Ovis canadensis canadensis isolate 43U chromosome 24 sequence : 36
Sus scrofa 18S ribosomal RNA gene, complete sequence : 35
Human DNA sequence from clone bP-2171C21 on chromosome 21, complete sequence : 35

SRR10168381, number of contigs 11914

Paenibacillus sordellii strain AM370 chromosome, complete genome : 196
Clostridiaceae bacterium 14S0207 chromosome, complete genome : 110
Beta vulgaris subsp. vulgaris cultivar KWS2320 chloroplast, complete genome : 74
Edwardsiella tarda strain KC-Pc-HB1 chromosome, complete genome : 64
Flavobacterium sp. xlx-214 chromosome, complete genome : 59
Neobacillus thermoproprius strain DUT50_236 chromosome, complete genome : 56
Beta vulgaris chloroplast sequence : 55
Plesiomonas shigelloides strain NCTC10360 genome assembly, chromosome: 1 : 52
Plesiomonas shigelloides strain MS-17-188 chromosome, complete genome : 52
Romboutsia ilealis strain CRIB genome assembly, chromosome: chr1 : 51

SRR10168382, number of contigs 1683

Pseudolabrys sp. FHR47 chromosome, complete genome : 857

Lutra lutra genome assembly, chromosome: 16 : 33

Mus musculus clone contig 1 chromocenter region genomic sequence : 29

Staphylococcus aureus strain WH9628 chromosome : 29

Sus scrofa 18S ribosomal RNA gene, complete sequence : 29

Human DNA sequence from clone bP-2171C21 on chromosome 21, complete sequence : 29

Homo sapiens clone N29M24 sequence : 29

Homo sapiens clone LA15_25H3 sequence : 29

Homo sapiens clone LA14_101B3 sequence : 29

Homo sapiens clone LA13_165F6 sequence : 29

SRR10168383, number of contigs 3991

Beta vulgaris subsp. *vulgaris* cultivar KWS2320 chloroplast, complete genome : 97

Beta vulgaris chloroplast sequence : 72

Eukaryotic synthetic construct chromosome 17 : 17

Homo sapiens DNA, chromosome 17, nearly complete genome : 16

Beta vulgaris subsp. *vulgaris* mitochondrial DNA, complete genome : 13

Manis javanica isolate MP_PG03-UM mitochondrion, complete genome : 12

Manis javanica isolate EP2 mitochondrion, complete genome : 12

Manis javanica isolate T298 mitochondrion, complete genome : 12

Manis javanica isolate MJA633 mitochondrion, complete genome : 11

Myroides phaeus strain 18QD1AZ29W chromosome, complete genome : 11

SRR10168384, number of contigs 11723

Beta vulgaris subsp. *vulgaris* cultivar KWS2320 chloroplast, complete genome : 74

Beta vulgaris chloroplast sequence : 53

Lutra lutra genome assembly, chromosome: 16 : 26

Mus musculus clone contig 1 chromocenter region genomic sequence : 21

Staphylococcus aureus strain WH9628 chromosome : 21

Sus scrofa 18S ribosomal RNA gene, complete sequence : 21

Human DNA sequence from clone bP-2171C21 on chromosome 21, complete sequence : 21

Homo sapiens clone N29M24 sequence : 21

Homo sapiens clone LA15_25H3 sequence : 21

Homo sapiens clone LA14_101B3 sequence : 21

SRR10168385, number of contigs 3661

Beta vulgaris subsp. *vulgaris* cultivar KWS2320 chloroplast, complete genome : 94

Beta vulgaris chloroplast sequence : 71

Acinetobacter bereziniae strain GD0320 chromosome, complete genome : 32

Acinetobacter bereziniae strain GD03185 chromosome, complete genome : 21
 Clostridiaceae bacterium 14S0207 chromosome, complete genome : 19
 Beta macrocarpa mitochondrion, complete genome : 18
 Acinetobacter bereziniae strain XH901, complete genome : 17
 Beta vulgaris subsp. maritima genotype male-fertile B mitochondrion, complete genome : 17
 Beta vulgaris subsp. maritima genotype male-fertile A mitochondrion, complete genome : 16
 Beta vulgaris subsp. vulgaris mitochondrial DNA, complete genome : 15

SRR10168386, number of contigs 1870

Manis javanica isolate EP2 mitochondrion, complete genome : 24
 Manis javanica isolate T298 mitochondrion, complete genome : 24
 Manis javanica isolate MJA633 mitochondrion, complete genome : 21
 Manis javanica isolate MP_PG03-UM mitochondrion, complete genome : 13
 Homo sapiens DNA, sequence_id: unplaced_0471 : 13
 Paenibacillus sordellii strain AM370 chromosome, complete genome : 12
 Manis pentadactyla mitochondrion, complete genome : 9
 Clostridium isatidis strain DSM 15098 chromosome, complete genome : 8
 Homo sapiens DNA, sequence_id: unplaced_0335 : 7
 Eukaryotic synthetic construct chromosome 15 : 6

SRR10168387, number of contigs 7626

Beta vulgaris subsp. vulgaris cultivar KWS2320 chloroplast, complete genome : 77
 Beta vulgaris chloroplast sequence : 56
 Lutra lutra genome assembly, chromosome: 16 : 19
 Staphylococcus aureus strain WH9628 chromosome : 14
 Acinetobacter bereziniae strain GD0320 chromosome, complete genome : 13
 Mus musculus clone contig 1 chromocenter region genomic sequence : 12
 Eukaryotic synthetic construct chromosome 17 : 12
 Beta vulgaris subsp. maritima genotype male-fertile B mitochondrion, complete genome : 12
 Manis pentadactyla mitochondrion, complete genome : 12
 Sus scrofa 18S ribosomal RNA gene, complete sequence : 12

SRR10168388, number of contigs 3512

Lutra lutra genome assembly, chromosome: 16 : 42
 Mus musculus clone contig 1 chromocenter region genomic sequence : 39
 Staphylococcus aureus strain WH9628 chromosome : 39
 Human DNA sequence from clone bP-2171C21 on chromosome 21, complete sequence : 39
 Homo sapiens clone N29M24 sequence : 39
 Homo sapiens clone LA15_25H3 sequence : 39
 Homo sapiens clone LA14_101B3 sequence : 39

Homo sapiens clone LA13_165F6 sequence : 39
Mouse DNA sequence from clone RP23-81C12 on chromosome 17, complete sequence :
39
Ovis canadensis canadensis isolate 43U chromosome 24 sequence : 39

SRR10168389, number of contigs 3544
Clostridium baratii strain CDC51267 chromosome, complete genome : 57
Clostridium baratii str. Sullivan, complete genome : 51
Lutra lutra genome assembly, chromosome: 16 : 33
Staphylococcus aureus strain WH9628 chromosome : 29
Sus scrofa 18S ribosomal RNA gene, complete sequence : 27
Homo sapiens clone N29M24 sequence : 26
Homo sapiens clone LA14_101B3 sequence : 26
Mus musculus clone contig 1 chromocenter region genomic sequence : 25
Mouse DNA sequence from clone RP23-81C12 on chromosome 17, complete sequence :
25
Homo sapiens clone LA15_25H3 sequence : 24

SRR10168390, number of contigs 6993
Trypanosoma cruzi cruzi strain Sylvio X10/cl1 chromosome TcI7 sequence : 27
Oscheius tipulae isolate CEW1 chromosome V : 27
Lactobacillus gastricus strain LG045 chromosome, complete genome : 24
[Candida] glabrata strain BG2 chromosome M : 23
Lutra lutra genome assembly, chromosome: 16 : 22
Staphylococcus condimentii strain DSM 11674, complete genome : 20
Staphylococcus condimentii strain NCTC13827 genome assembly, chromosome: 1 : 20
Staphylococcus condimentii strain St0 2014-01, complete genome : 17
Lactobacillus mucosae LM1, complete genome : 17
Staphylococcus condimentii strain FDAARGOS_1148 chromosome, complete genome : 16

SRR10168391, number of contigs 10811
Myroides phaeus strain 18QD1AZ29W chromosome, complete genome : 337
Acinetobacter bereziniae strain GD0320 chromosome, complete genome : 168
Acinetobacter bereziniae strain GD03185 chromosome, complete genome : 148
Acinetobacter bereziniae strain XH901, complete genome : 145
Myroides odoratimimus strain G13 chromosome, complete genome : 117
Vagococcus carniphilus strain ATCC BAA-640 chromosome, complete genome : 112
Myroides sp. A21, complete genome : 100
Myroides odoratimimus strain PR63039, complete genome : 92
Beta vulgaris subsp. vulgaris cultivar KWS2320 chloroplast, complete genome :
91
Myroides sp. ZB35, complete genome : 86

SRR10168392, number of contigs 21005

Lutra lutra genome assembly, chromosome: 16 : 107
 Staphylococcus aureus strain WH9628 chromosome : 88
 Sus scrofa 18S ribosomal RNA gene, complete sequence : 87
 Mouse DNA sequence from clone RP23-81C12 on chromosome 17, complete sequence :
 82
 Homo sapiens clone LA14_101B3 sequence : 79
 Homo sapiens clone N29M24 sequence : 77
 Homo sapiens clone LA15_25H3 sequence : 77
 Mus musculus clone contig 1 chromocenter region genomic sequence : 76
 Homo sapiens clone LA13_165F6 sequence : 75
 Human DNA sequence from clone bP-2171C21 on chromosome 21, complete sequence :
 74

SRR10168393, number of contigs 10877
 Homo sapiens DNA, chromosome 16, nearly complete genome : 221
 Eukaryotic synthetic construct chromosome 16 : 213
 Homo sapiens 16p13.3 sequence section 1 of 8 : 98
 Homo sapiens interleukin 9 receptor (IL9Rps) pseudogene, complete sequence; and
 DNA-directed RNA polymerases III 12.5 kDa polypeptide (POLR3K), U11/U12 snRNP 25
 kDa protein (C16orf33), RHBDG1 (C16orf8), DNA-3-methyladenine glycosylase (MPG),
 -14 gene protein (C16orf35), hemoglobin zeta subunit (HBZ), HBD (HBD), alpha
 globin (HBA2), alpha globin (HBA1), theta globin (HBQ1), and putative RNA-
 binding protein Luc7-like 1 (LUC7L) genes, complete cds : 96
 Homo sapiens 16p13.3 sequence section 2 of 8 : 67
 Homo sapiens chromosome 11, clone CTD-2643I7, complete sequence : 66
 Eukaryotic synthetic construct chromosome 19 : 50
 Eukaryotic synthetic construct chromosome 17 : 47
 Homo sapiens DNA, chromosome 17, nearly complete genome : 46
 Homo sapiens DNA, chromosome 19, nearly complete genome : 44

[57]: *### Get specific contigs matching a species/name*

```
[58]: def write_contigs(sra, match_names):
    accessions, descriptions, counts, total=get_asc_descr_count(sra)
    asc_matches=[]
    for m in match_names:
        for a,d in zip(accessions, descriptions):
            if m.lower() in d.lower():
                asc_matches.append(a)
    gi_matches=[]
    for a in asc_matches:
        idx=ACCESSIONS.index(a)
        gi=GIS[idx]
        gi_matches.append(gi)
```



```

contigs=[]
path = BASE_PATH+sra+'/magic_blast/'
subset_f = open(path+f'{sra}_{match_names[0].replace(" ", "_")}_subset_{f_contigs_file_tail}', 'w')
with open(path+f'{sra}_{f_contigs_file_tail}', 'r') as f:
    lines = [line for line in f]
    for line in lines:
        for gi in gi_matches:
            if gi in line:
                parts=line.split('\t')
                idx=GIS.index(gi)
                asc=ACCESSIONS[idx]
                t=TITLES[idx]
                ps=parts[:2]
                pe=parts[3:]
                ps.append(asc+' '+t.rstrip('\n'))
                parts=ps+pe
                p='\t'.join(parts)
                subset_f.write(p)
subset_f.close()

```

```
[ ]:
```

```
[59]: #set_accessions()
      #assert ACCESSIONS is not None

```

```
[60]: #match_names=['vector']
      #for sra in sra_list:
      #    write_contigs(sra, match_names)

```

```
[61]: #match_names=['plasmid']
      #for sra in sra_list:
      #    write_contigs(sra, match_names)

```

```
[62]: #match_names=['mustela']
      #for sra in sra_list:
      #    write_contigs(sra, match_names)

```

```
[63]: #match_names=['virus']
      #for sra in sra_list:
      #    write_contigs(sra, match_names)

```